Study guide: Introduction to finite element methods

Hans Petter Langtangen 1,2

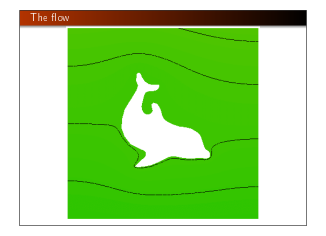
Center for Biomedical Computing, Simula Research Laboratory¹

Department of Informatics, University of Oslo²

Oct 16, 2014

Can with ease solve PDEs in domains with complex geometry Can with ease provide higher-order approximations Has (in simpler stationary problems) a rigorous mathematical analysis framework (not much considered here)

Domain for flow around a dolphin



Stationary PDEs: Transform the PDE problem to a variational form Define function approximation over finite elements Use a machinery to derive linear systems Solve linear systems Time-dependent PDEs: Finite elements in space Finite difference (or ODE solver) in time

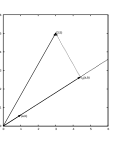
We start with function approximation, then we treat PDEs

Learning strategy

Start with approximation of functions, not PDEs
Introduce finite element approximations
See later how this machinery is applied to PDEs

Reason:
The finite element method has many concepts and a jungle of details. This learning strategy minimizes the mixing of ideas, concepts, and technical details.

Approximation in vector spaces



Approximation set-up

General idea of finding an approximation u(x) to some given f(x):

$$u(x) = \sum_{i=0}^{N} c_i \psi_i(x)$$

wh ere

- \bullet $\psi_i(x)$ are prescribed functions
- ullet $c_{i},\ i=0,\ldots,N$ are unknown coefficients to be determined

How to determine the coefficients?

We shall address three approaches:

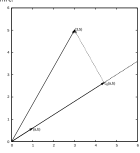
- The least squares method
- The projection (or Galerkin) method
- The interpolation (or collocation) method

Underlying motivation for our notation

Our mathematical framework for doing this is phrased in a way such that it becomes easy to understand and use the FEniCS software package for finite element computing.

Approximation of planar vectors; problem

Given a vector f = (3, 5), find an approximation to f directed along a given line.



Approximation of planar vectors; vector space terminology

$$V=\operatorname{\mathsf{span}}\left\{oldsymbol{\psi}_0
ight\}$$

- ullet ψ_0 is a basis vector in the space V
- Seek $u = c_0 \psi_0 \in V$
- ullet Determine c_0 such that $oldsymbol{u}$ is the "best" approximation to $oldsymbol{f}$
- Visually, "best" is obvious

Define

- the error e = f u
- ullet the (Eucledian) scalar product of two vectors: (u,v)
- the norm of e: $||e|| = \sqrt{(e,e)}$

The least squares method; principle

- Idea: find c_0 such that ||e|| is minimized
- Mathematical convenience: minimize $E = ||e||^2$

$$\frac{\partial E}{\partial c_0} = 0$$

The least squares method; calculations

$$E(c_0) = (e, e) = (f - u, (f - u)) = (f - c_0\psi_0, f - c_0\psi_0)$$

= $(f, f) - 2c_0(f, \psi_0) + c_0^2(\psi_0, \psi_0)$

$$\frac{\partial E}{\partial c_0} = -2(\mathbf{f}, \psi_0) + 2c_0(\psi_0, \psi_0) = 0 \tag{1}$$

$$c_0 = \frac{(f, \psi_0)}{(\psi_0, \psi_0)} = \frac{3a + 5b}{a^2 + b^2}$$

Observation to be used later: the vanishing derivative (1) can be alternatively written as $\frac{1}{2}$

$$(e, \psi_0) = 0$$

The projection (or Galerkin) method

- Last slide: min E is equivalent with $(e, \psi_0) = 0$
- $(e, \psi_0) = 0$ implies $(e, \mathbf{v}) = 0$ for any $\mathbf{v} \in V$
- That is: instead of using the least-squares principle, we can require that e is orthogonal to any v ∈ V (visually clear, but can easily be computed too)
- Precise math: find c_0 such that (e, v) = 0, $\forall v \in V$
- ullet Equivalent (see notes): find c_0 such that $(oldsymbol{e}, oldsymbol{\psi}_0) = 0$
- Insert $e = f c_0 \psi_0$ and solve for c_0
- Same equation for c_0 and hence same solution as in the least squares method

Approximation of general vectors

Given a vector f, find an approximation $u \in V$:

$$V = \operatorname{span} \{\psi_0, \dots, \psi_N\}$$

We have a set of linearly independent basis vectors ψ_0,\dots,ψ_N . Any $u\in V$ can then be written as

$$u = \sum_{i=0}^{N} c_j \psi_j$$

The least squares method

Idea: find c_0, \ldots, c_N such that $E = ||e||^2$ is minimized, e = f - u.

$$E(c_0, ..., c_N) = (e, e) = (f - \sum_j c_j \psi_j, f - \sum_j c_j \psi_j)$$

= $(f, f) - 2 \sum_{j=0}^{N} c_j (f, \psi_j) + \sum_{p=0}^{N} \sum_{q=0}^{N} c_p c_q (\psi_p, \psi_q)$

$$\frac{\partial E}{\partial c_i} = 0, \quad i = 0, \dots, N$$

After some work we end up with a linear system

$$\sum_{j=0}^{N} A_{i,j} c_j = b_i, \quad i = 0, \dots, N$$
 (2)

$$A_{i,j} = (\psi_i, \psi_j) \tag{3}$$

The projection (or Galerkin) method

Can be shown that minimizing ||e|| implies that e is orthogonal to all $\mathbf{v} \in V$:

$$(e, \mathbf{v}) = 0, \quad \forall \mathbf{v} \in V$$

which implies that e most be orthogonal to each basis vector:

$$(e, \psi_i) = 0, \quad i = 0, ..., N$$

This orthogonality condition is the principle of the projection (or Galerkin) method. Leads to the same linear system as in the least squares method.

Approximation of functions

Let V be a function space spanned by a set of basis functions ψ_0, \dots, ψ_N ,

$$V = \operatorname{span} \{\psi_0, \dots, \psi_N\}$$

Find $u \in V$ as a linear combination of the basis functions:

$$u = \sum_{j \in \mathcal{I}_{\sigma}} c_j \psi_j, \quad \mathcal{I}_{\sigma} = \{0, 1, \dots, N\}$$

The least squares method can be extended from vectors to functions

As in the vector case, minimize the (square) norm of the error, E, with respect to the coefficients $c_i, j \in \mathcal{I}_s$:

$$E = (e, e) = (f - u, f - u) = (f(x) - \sum_{j \in \mathcal{I}_s} c_j \psi_j(x), f(x) - \sum_{j \in \mathcal{I}_s} c_j \psi_j(x))$$

$$\frac{\partial E}{\partial c_i} = 0, \quad i = \in \mathcal{I}_s$$

But what is the scalar product when ψ_i is a function?

$$(f,g) = \int_{\Omega} f(x)g(x) \, dx$$

(natural extension from Eucledian product $(u, v) = \sum_i u_i v_i$)

The least squares method; details

$$E(c_0, ..., c_N) = (e, e) = (f - u, f - u)$$

= $(f, f) - 2 \sum_{j \in \mathcal{I}_s} c_j(f, \psi_i) + \sum_{\rho \in \mathcal{I}_s} \sum_{q \in \mathcal{I}_s} c_\rho c_q(\psi_\rho, \psi_q)$

$$\frac{\partial E}{\partial c_i} = 0, \quad i = \in \mathcal{I}_s$$

The computations are identical to the vector case, and consequently we get a linear system

$$\sum_{i\in\mathcal{I}_s}^N A_{i,j}c_j = b_i, \ i\in\mathcal{I}_s, \quad A_{i,j} = (\psi_i,\psi_j), \ b_i = (f,\psi_i)$$

The projection (or Galerkin) method

As before, minimizing (e, e) is equivalent to

$$(e, \psi_i) = 0, \quad i \in \mathcal{I}_s$$

which is equivalent to

$$(e, v) = 0, \forall v \in V$$

which is the projection (or Galerkin) method

The algebra is the same as in the multi-dimensional vector case, and we get the same linear system as arose from the least squares method.

Example: linear approximation; problem

Problem

Approximate a parabola $f(x) = 10(x-1)^2 - 1$ by a straight line.

$$V = \operatorname{span} \{1, x\}$$

That is, $\psi_0(x)=1$, $\psi_1(x)=x$, and N=1. We seek

$$u = c_0\psi_0(x) + c_1\psi_1(x) = c_0 + c_1x$$

Example: linear approximation; solution

$$A_{0,0} = (\psi_0, \psi_0) = \int_1^2 1 \cdot 1 \, dx = 1$$

$$A_{0,1} = (\psi_0, \psi_1) = \int_1^2 1 \cdot x \, dx = 3/2$$

$$A_{1,0} = A_{0,1} = 3/2$$

$$A_{1,1} = (\psi_1, \psi_1) = \int_1^2 x \cdot x \, dx = 7/3$$

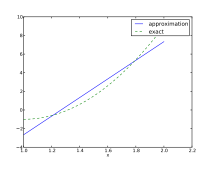
$$b_1 = (f, \psi_0) = \int_1^2 (10(x - 1)^2 - 1) \cdot 1 \, dx = 7/3$$

$$b_2 = (f, \psi_1) = \int_1^2 (10(x - 1)^2 - 1) \cdot x \, dx = 13/3$$

Solution of 2x2 linear system:

$$c_0 = -38/3$$
, $c_1 = 10$, $u(x) = 10x - \frac{38}{3}$

Example: linear approximation; plot



Implementation of the least squares method; ideas

Consider symbolic computation of the linear system, where

- f(x) is given as a sympy expression f (involving the symbol x),
- ullet psi is a list of $\{\psi_i\}_{i\in\mathcal{I}_s}$
- ullet Omega is a 2-tuple/list holding the domain Ω

Carry out the integrations, solve the linear system, and return $u(x) = \sum_i c_i \psi_i(x)$

Implementation of the least squares method; symbolic code

Observe: symmetric coefficient matrix so we can halve the integrations.

Improved code if symbolic integration fails

- If sp.integrate fails, it returns an sp.Integral object. We can test on this object and fall back on numerical integration.
- We can include a boolean argument symbolic to explicitly choose between symbolic and numerical computing.

Implementation of the least squares method; plotting

```
def comparison_plot(f, u, Omega, filename='tmp.pdf'):
    x = sp.Symbol('x')
# Turn f and u to ordinary Python functions
    f = sp.lambdify([x], f, modules="numpy")
    u = sp.lambdify([x], u, modules="numpy")
    resolution = 401 # no of points in plot
    xcoor = linspace(Omega[0], Omega[1], resolution)
    exact = f(xcoor)
    approx = u(xcoor)
    plot(xcoor, approx)
```

plot(xcoor, exact)
legend(['approximation', 'exact'])
savefig(filename)

All code in module approx 1D.py

Compare f and u visually:

Perfect approximation; parabola approximating parabola

- What if we add $\psi_2 = x^2$ to the space V?
- That is, approximating a parabola by any parabola?
- (Hopefully we get the exact parabola!)

```
>>> from approx1D import *
>>> x = sp.Symbol('x')
>>> f = 10*(x-1)**2-1
>>> u, c = least squares(f=f, psi=[1, x, x**2], Omega=[1, 2])
>>> print u
10**x**2 - 20*x + 9
>>> print sp.expand(f)
10*x**2 - 20*x + 9
```

Perfect approximation; the general result

- What if we use $\psi_i(x) = x^i$ for i = 0, ..., N = 40?
- The output from least_squares is $c_i = 0$ for i > 2

General result

If $f \in V$, least squares and projection/Galerkin give u = f.

Perfect approximation; proof of the general result

If $f \in V$, $f = \sum_{j \in \mathcal{I}_s} d_j \psi_j$, for some $\{d_i\}_{i \in \mathcal{I}_s}$. Then

$$b_i = (f, \psi_i) = \sum_{j \in \mathcal{I}_s} d_j(\psi_j, \psi_i) = \sum_{j \in \mathcal{I}_s} d_j A_{i,j}$$

The linear system $\sum_j A_{i,j} c_j = b_i, \ i \in \mathcal{I}_s$, is then

$$\sum_{j \in \mathcal{I}_s} c_j A_{i,j} = \sum_{j \in \mathcal{I}_s} d_j A_{i,j}, \quad i \in \mathcal{I}_s$$

which implies that $c_i = d_i$ for $i \in \mathcal{I}_s$ and u is identical to f.

Finite-precision/numerical computations; question

The previous computations were symbolic. What if we solve the linear system numerically with standard arrays?

That is, f is parabola, but we approximate with

$$u(x) = c_0 + c_1 x + c_2 x^2 + c_3 x^3 + \dots + c_N x^N$$

We expect $c_2 = c_3 = \cdots = c_N = 0$ since $f \in V$ implies u = f.

Will we get this result with finite precision computer arithmetic?

Finite-precision/numerical computations; results

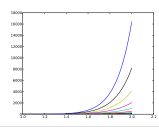
exact	sympy	numpy 32	numpy 64
9	9.62	5.57	8.98
-20	-23.39	-7.65	-19.93
10	17.74	-4.50	9.96
0	-9.19	4.13	-0.26
0	5.25	2.99	0.72
0	0.18	-1.21	-0.93
0	-2.48	-0.41	0.73
0	1.81	-0.013	-0.36
0	-0.66	0.08	0.11
0	0.12	0.04	-0.02
0	-0.001	-0.02	0.002

- Column 2: matrix and lu_solve from sympy.mpmath.fp
- Column 3: numpy matrix with 4-byte floats
- Column 4: numpy matrix with 8-byte floats

The ill-conditioning is due to almost linearly dependent basis functions for large $\,N\,$

- Significant round-off errors in the numerical computations (!)
- But if we plot the approximations they look good (!)

Source or problem: the \mathbf{x}^i functions become almost linearly dependent as i grows:



Ill-conditioning: general conclusions

- Almost linearly dependent basis functions give almost singular matrices
- Such matrices are said to be *ill conditioned*, and Gaussian elimination is severely affected by round-off errors
- The basis $1, x, x^2, x^3, x^4, \dots$ is a bad basis
- ullet Polynomials are fine as basis, but the more orthogonal they are, $(\psi_i,\psi_i)pprox 0$, the better

Fourier series approximation; problem and code

Let's approximate f by a typical Fourier series expansion

$$u(x) = \sum_i a_i \sin i\pi x = \sum_{j=0}^N c_j \sin((j+1)\pi x)$$

which means that

$$V = \operatorname{span} \left\{ \sin \pi x, \sin 2\pi x, \dots, \sin (N+1)\pi x \right\}$$

Computations using the least_squares function:

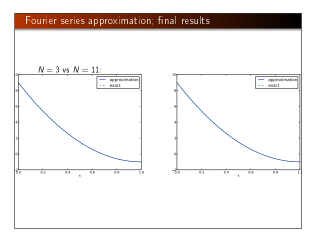
```
N = 3
from sympy import sin, pi
psi = [sin(pi*(i+1)*x) for i in range(N+1)]
f = 10*(x-1)**2 - 1
Omega = [0, 1]
u, c = least_squares(f, psi, Omega)
comparison_plot(f, u, Omega)
```


Fourier series approximation; improvements

- \bullet Considerably improvement with ${\it N}=11$, but still undesired discrepancy at x=0 and x=1
- Possible remedy: add a term that leads to correct boundary values

$$u(x) = f(0)(1-x) + xf(1) + \sum_{j \in \mathcal{I}_s} c_j \psi_j(x)$$

The extra terms ensure u(0)=f(0) and u(1)=f(1) and is a strikingly good help to get a good approximation!



Orthogonal basis functions

This choice of sine functions as basis functions is popular because

- the basis functions are orthogonal: $(\psi_i, \psi_i) = 0$
- implying that $A_{i,j}$ is a diagonal matrix
- implying that we can solve for $c_i = 2 \int_0^1 f(x) \sin((i+1)\pi x) dx$
- ullet and what we get is the standard Fourier sine series of f

In general, for an orthogonal basis, $A_{i,j}$ is diagonal and we can easily solve for c_i :

$$c_i = \frac{b_i}{A_{i,i}} = \frac{(f, \psi_i)}{(\psi_i, \psi_i)}$$

Function for the least squares method with orthogonal basis functions

```
def least_squares_orth(f, psi, Omega):
    N = len(psi) - 1
    A = [0] *(N+1)
    b = [0] *(N+1)
    x = sp. Symbol('x')
    for i in range(N+1):
        A [i] = sp. integrate(psi[i] **2, (x, Omega[0], Omega[1]))
        b[i] = sp. integrate(psi[i] *f, (x, Omega[0], Omega[1]))
    c = [b[i]/A[i] for i in range(len(b))]
    u = 0
    for i in range(len(psi)):
        u + c c[i] *psi[i]
    return u, c
```

Function for the least squares method with orthogonal basis functions; symbolic and numerical integration

Extensions:

- We can choose between symbolic or numerical integration (symbolic argument).
- If symbolic, we fall back on numerical integration after failure (sp.Integral is returned from sp.integrate).

The collocation or interpolation method; implementation

points holds the interpolation/collocation points

Lagrange polynomials; motivation and ideas

Motivation

- The interpolation/collocation method avoids integration
- ullet With a diagonal matrix $A_{i,j}=\psi_j(x_i)$ we can solve the linear system by hand

The Lagrange interpolating polynomials ψ_j have the property that

$$\psi_i(x_j) = \delta_{ij}, \quad \delta_{ij} = \begin{cases} 1, & i = j \\ 0, & i \neq j \end{cases}$$

Hence, $c_i = f(x_i)$ and

$$u(x) = \sum_{j \in \mathcal{I}_s} f(x_i) \psi_i(x)$$

 Lagrange polynomials and interpolation/collocation look convenient

The collocation or interpolation method; ideas and math

Here is another idea for approximating f(x) by $u(x) = \sum_i c_i \psi_i$:

- Force $u(x_i) = f(x_i)$ at some selected *collocation* points $\{x_i\}_{i \in \mathcal{T}_*}$
- Then u is said to interpolate f
- The method is known as interpolation or collocation

$$u(x_i) = \sum_{j \in \mathcal{I}_s} c_j \psi_j(x_i) = f(x_i) \quad i \in \mathcal{I}_s, N$$

This is a linear system with no need for integration:

$$\sum_{j\in\mathcal{I}_s} A_{i,j}c_j = b_i, \quad i\in\mathcal{I}_s$$
 (5)

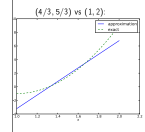
$$A_{i,j} = \psi_j(x_i) \tag{6}$$

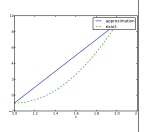
$$b_i = f(x_i) \tag{7}$$

No symmetric matrix: $\psi_i(x_i) \neq \psi_i(x_i)$ in general

The collocation or interpolation method; approximating a parabola by linear functions

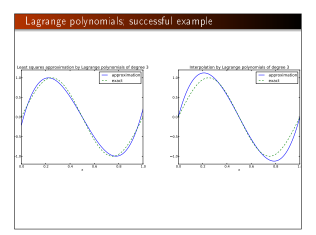
- Potential difficulty: how to choose x_i ?
- The results are sensitive to the points!

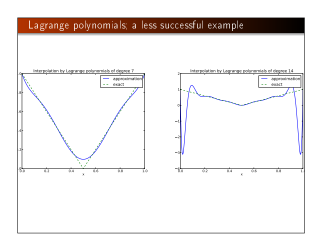


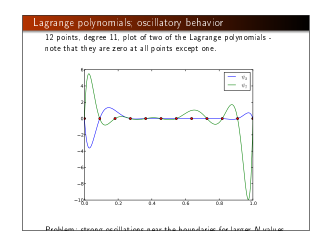


Lagrange polynomials; formula and code

$$\begin{split} \psi_i(x) &= \prod_{j=0, j \neq i}^N \frac{x-x_j}{x_i-x_j} = \frac{x-x_0}{x_i-x_0} \cdots \frac{x-x_{i-1}}{x_i-x_{i-1}} \frac{x-x_{i+1}}{x_i-x_{i+1}} \cdots \frac{x-x_N}{x_i-x_N} \\ &\text{def Lagrange_polynomial}(x, i, points): \\ &\text{p = 1} \\ &\text{for k in range}(len(points)): \\ &\text{if k != i:} \\ &\text{p *= (x - points[k]) / (points[i] - points[k])} \\ &\text{return p} \end{split}$$





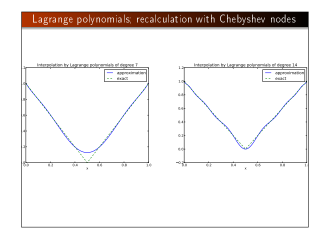


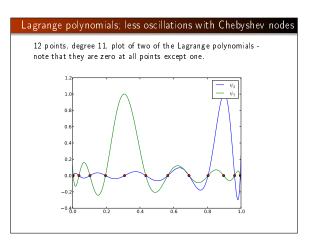
Lagrange polynomials; remedy for strong oscillations

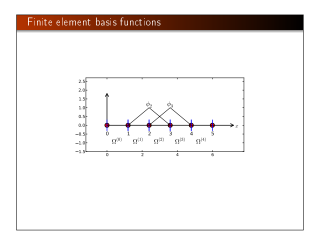
The oscillations can be reduced by a more clever choice of interpolation points, called the *Chebyshev nodes*:

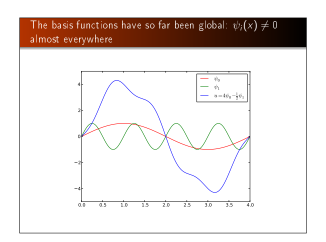
$$x_i = \frac{1}{2}(a+b) + \frac{1}{2}(b-a)\cos\left(\frac{2i+1}{2(N+1)}\rho i\right), \quad i = 0..., N$$

on an interval [a, b].



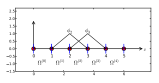


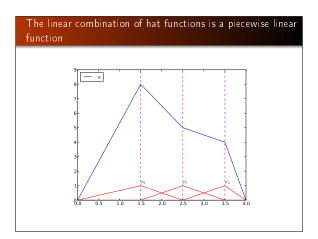




In the finite element method we use basis functions with local support

- Local support: $\psi_i(x) \neq 0$ for x in a small subdomain of Ω
- Typically hat-shaped
- u(x) based on these ψ_i is a piecewise polynomial defined over many (small) subdomains
- We introduce φ_i as the name of these finite element hat functions (and for now choose $\psi_i = \varphi_i$)





Elements and nodes

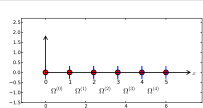
Split Ω into non-overlapping subdomains called *elements*:

$$\Omega = \Omega^{(0)} \cup \cdots \cup \Omega^{(N_e)}$$

On each element, introduce points called $nodes: x_0, \ldots, x_{N_n}$

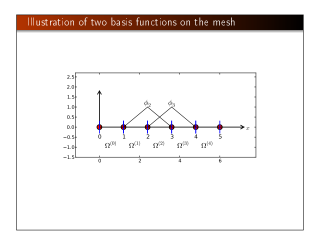
- The finite element basis functions are named $\varphi_i(x)$
- $ullet arphi_i = 1$ at node i and 0 at all other nodes
- ullet $arphi_i$ is a Lagrange polynomial on each element
- ullet For nodes at the boundary between two elements, $arphi_i$ is made up of a Lagrange polynomial over each element

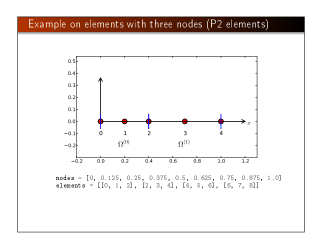
Example on elements with two nodes (P1 elements)

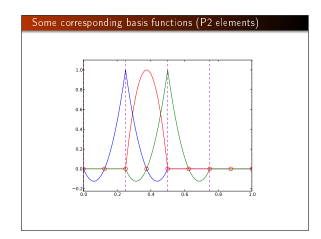


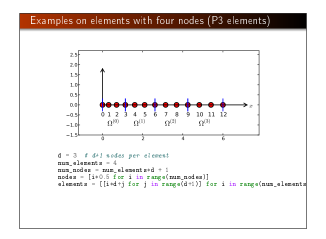
Data structure: nodes holds coordinates or nodes, elements holds the node numbers in each element

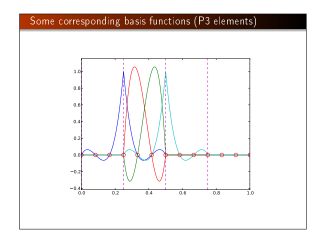
nodes = [0, 1.2, 2.4, 3.6, 4.8, 5] elements = [[0, 1], [1, 2], [2, 3], [3, 4], [4, 5]]

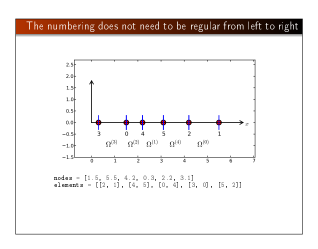












Interpretation of the coefficients c_i

Important property: c_i is the value of u at node i, x_i :

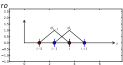
$$u(x_i) = \sum_{j \in \mathcal{I}_s} c_j \varphi_j(x_i) = c_i \varphi_i(x_i) = c_i$$

because $\varphi_i(x_i) = 0$ if $i \neq j$ and $\varphi_i(x_i) = 1$

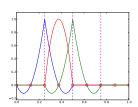
Properties of the basis functions

- $\varphi_i(x) \neq 0$ only on those elements that contain global node i
- $\varphi_i(x)\varphi_j(x) \neq 0$ if and only if i and j are global node numbers in the same element

Since $A_{i,j} = \int \varphi_i \varphi_j \, \mathrm{d}x$, most of the elements in the coefficient matrix will be zero____

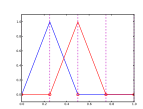


How to construct quadratic φ_i (P2 elements)



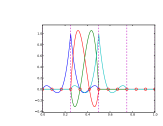
- Associate Lagrange polynomials with the nodes in an element
- When the polynomial is 1 on the element boundary, combine it with the polynomial in the neighboring element that is also 1 at the same point

Example on linear φ_i (P1 elements)

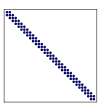


$$\varphi_i(x) = \begin{cases} 0, & x < x_{i-1} \\ (x - x_{i-1})/h & x_{i-1} \le x < x_i \\ 1 - (x - x_i)/h, & x_i \le x < x_{i+1} \\ 0, & x \ge x_{i+1} \end{cases}$$

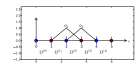
Example on cubic φ_i (P3 elements)



Calculating the linear system for c_i



Computing a specific matrix entry (1)

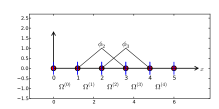


 $A_{2,3}=\int_{\Omega} arphi_2 arphi_3 dx\colon arphi_2 arphi_3
eq 0$ only over element 2. There,

$$\varphi_3(x) = (x - x_2)/h, \quad \varphi_2(x) = 1 - (x - x_2)/h$$

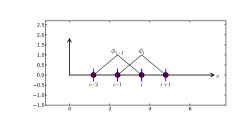
$$A_{2,3} = \int_{\Omega} \varphi_2 \varphi_3 \, \mathrm{d}x = \int_{x_2}^{x_3} \left(1 - \frac{x - x_2}{h} \right) \frac{x - x_2}{h} \, \mathrm{d}x = \frac{h}{6}$$

Computing a specific matrix entry (2)



$$A_{2,2} = \int_{x_1}^{x_2} \left(\frac{x - x_1}{h} \right)^2 dx + \int_{x_2}^{x_3} \left(1 - \frac{x - x_2}{h} \right)^2 dx = \frac{2h}{3}$$

Calculating a general row in the matrix; figure



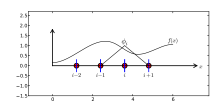
$$A_{i,i-1} = \int_{\Omega} \varphi_i \varphi_{i-1} \, \mathrm{d}x = ?$$

Calculating a general row in the matrix; details

$$\begin{split} A_{i,i-1} &= \int_{\Omega} \varphi_i \varphi_{i-1} \, \mathrm{d} x \\ &= \underbrace{\int_{x_{i-1}}^{x_{i-1}} \varphi_i \varphi_{i-1} \, \mathrm{d} x}_{\varphi_i = 0} + \underbrace{\int_{x_{i-1}}^{x_i} \varphi_i \varphi_{i-1} \, \mathrm{d} x}_{\varphi_{i-1}} + \underbrace{\int_{x_i}^{x_{i+1}} \varphi_i \varphi_{i-1} \, \mathrm{d} x}_{\varphi_{i-1} = 0} \\ &= \underbrace{\int_{x_{i-1}}^{x_i} \underbrace{\left(\frac{x - x_i}{h}\right)}_{\varphi_i(x)} \underbrace{\left(1 - \frac{x - x_{i-1}}{h}\right)}_{\varphi_{i-1}(x)} \, \mathrm{d} x = \frac{h}{6} \end{split}$$

- $A_{i,i+1} = A_{i,i-1}$ due to symmetry
- $A_{i,i} = 2h/3$ (same calculation as for $A_{2,2}$)
- $A_{0.0} = A_{N.N} = h/3$ (only one element)

Calculation of the right-hand side



$$b_{i} = \int_{\Omega} \varphi_{i}(x) f(x) dx = \int_{x_{i-1}}^{x_{i}} \frac{x - x_{i-1}}{h} f(x) dx + \int_{x_{i}}^{x_{i+1}} \left(1 - \frac{x - x_{i}}{h}\right) f(x)$$

Need a specific f(x) to do more...

Specific example with two elements; linear system and solution

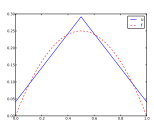
- f(x) = x(1-x) on $\Omega = [0,1]$
- ullet Two equal-sized elements [0,0.5] and [0.5,1]

$$A = \frac{h}{6} \begin{pmatrix} 2 & 1 & 0 \\ 1 & 4 & 1 \\ 0 & 1 & 2 \end{pmatrix}, \quad b = \frac{h^2}{12} \begin{pmatrix} 2 - 3h \\ 12 - 14h \\ 10 - 17h \end{pmatrix}$$

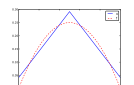
$$c_0 = \frac{h^2}{6}$$
, $c_1 = h - \frac{5}{6}h^2$, $c_2 = 2h - \frac{23}{6}h^2$

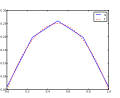
Specific example with two elements; plot

$$u(x) = c_0\varphi_0(x) + c_1\varphi_1(x) + c_2\varphi_2(x)$$



Specific example with four elements; plot

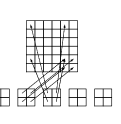




Specific example: what about P2 elements?

Recall: if $f \in V$, u becomes exact. When f is a parabola, any choice of P2 elements (1 or many) will give u = fexactly. The same is true for P3, P4, ... elements since all of them can represent a 2nd-degree polynomial exactly.

Assembly of elementwise computations

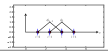


Split the integrals into elementwise integrals

$$A_{i,j} = \int_{\Omega} \varphi_i \varphi_j dx = \sum_{e} \int_{\Omega(e)} \varphi_i \varphi_j dx, \quad A_{i,j}^{(e)} = \int_{\Omega(e)} \varphi_i \varphi_j dx$$

Important observations:

- $A_{i,j}^{(e)} \neq 0$ if and only if i and j are nodes in element e (otherwise no overlap between the basis functions)
 All the nonzero elements in $A_{i,j}^{(e)}$ are collected in an element
- ullet The element matrix has contributions from the $arphi_i$ functions associated with the nodes in element
- It is convenient to introduce a local numbering of the nodes in an element: $0, 1, \ldots, d$

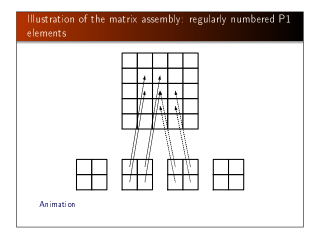


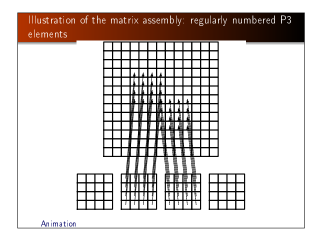
The element matrix and local vs global node numbers

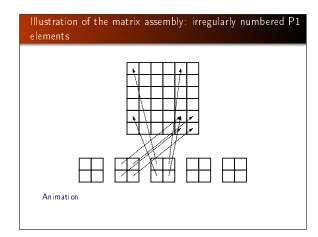
$$\tilde{A}^{(e)} = \{\tilde{A}_{r,s}^{(e)}\}, \quad \tilde{A}_{r,s}^{(e)} = \int_{\Omega^{(e)}} \varphi_{q(e,r)} \varphi_{q(e,s)} dx, \quad r, s \in I_d = \{0,\ldots,d\}$$

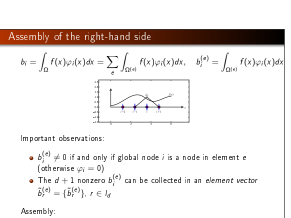
- r, s run over local node numbers in an element: $0, 1, \ldots, d$
- i,j run over global node numbers $i,j \in \mathcal{I}_s = \{0,1,\ldots,N\}$
- i = q(e, r): mapping of local node number r in element e to the global node number i (math equivalent to i=elements[e][r])
- Add $\tilde{A}_{r,s}^{(e)}$ into the global $A_{i,i}$ (assembly)

$$A_{q(e,r),q(e,s)} := A_{q(e,r),q(e,s)} + \tilde{A}_{r,s}^{(e)}, \quad r,s \in I_d$$









 $b_{q(e,r)} := b_{q(e,r)} + \tilde{b}_{r}^{(e)}, \quad r, s \in I_{d}$

Mapping to a reference element $\vec{A}_{r,s}^{(e)} = \int_{\Omega^{(e)}} \varphi_{q(e,r)}(x) \varphi_{q(e,s)}(x) dx = \int_{x_L}^{x_R} \varphi_{q(e,r)}(x) \varphi_{q(e,s)}(x) dx$ we now map $[x_L, x_R]$ to a standardized reference element domain [-1,1] with local coordinate X

We use affine mapping: linear stretch of
$$X\in[-1,1]$$
 to $x\in[x_L,x_R]$
$$x=\frac{1}{2}(x_L+x_R)+\frac{1}{2}(x_R-x_L)X$$
 or rewritten as
$$x=x_m+\frac{1}{2}hX, \qquad x_m=(x_L+x_R)/2, \quad h=x_R-x_L$$

Integral transformation

Reference element integration: just change integration variable from x to X. Introduce local basis function

$$\tilde{\varphi}_r(X) = \varphi_{q(e,r)}(x(X))$$

$$\tilde{A}_{r,s}^{(e)} = \int_{\Omega^{(e)}} \varphi_{q(e,r)}(x) \varphi_{q(e,s)}(x) dx = \int_{-1}^{1} \tilde{\varphi}_{r}(X) \tilde{\varphi}_{s}(X) \underbrace{\frac{dx}{dX}}_{\det J = h/2} dX = \int_{-1}^{1} \tilde{\varphi}_{r}(X) \tilde{\varphi}_{s}(X) \underbrace{\frac{dx}{dX}}_{\det J = h/2} dX = \int_{-1}^{1} \tilde{\varphi}_{r}(X) \tilde{\varphi}_{s}(X) \underbrace{\frac{dx}{dX}}_{\det J = h/2} dX = \underbrace{\int_{-1}^{1} \tilde{\varphi}_{r}(X) \tilde{\varphi}_{s}(X)}_{\det J = h/2} dX = \underbrace{\int_{-1}^{1} \tilde{\varphi}_{r}(X)}_{\det J}_{\det J$$

$$\tilde{b}_r^{(e)} = \int_{\Omega^{(e)}} f(x) \varphi_{q(e,r)}(x) dx = \int_{-1}^1 f(x(X)) \tilde{\varphi}_r(X) \det J \, dX$$

Advantages of the reference element

- Always the same domain for integration: [-1, 1]
- ullet We only need formulas for $ilde{arphi}_r(X)$ over one element (no piecewise polynomial definition)
- \bullet $\tilde{\varphi}_r(X)$ is the same for all elements: no dependence on element length and location, which is "factored out" in the mapping and $\det J$

Standardized basis functions for P1 elements

$$\tilde{\varphi}_0(X) = \frac{1}{2}(1 - X) \tag{8}$$

$$\tilde{\varphi}_1(X) = \frac{1}{2}(1+X) \tag{9}$$

Note: simple polynomial expressions (no need to consider piecewisely defined functions)

Standardized basis functions for P2 elements

$$\tilde{\varphi}_0(X) = \frac{1}{2}(X - 1)X$$
(10)
$$\tilde{\varphi}_1(X) = 1 - X^2$$
(11)

$$\tilde{\rho}_1(X) = 1 - X^2 \tag{11}$$

$$\tilde{\varphi}_2(X) = \frac{1}{2}(X+1)X$$
 (12)

Easy to generalize to arbitrary order!

How to find the polynomial expressions?

Three alternatives:

- Map the global basis function $\varphi_i(x)$ over an element to X
- ② Compute $\tilde{\varphi}_r(X)$ from scratch using
 - a given polynomial order d
 - $\tilde{\varphi}_r(X) = 1 \text{ at local node } 1$
 - $\tilde{\varphi}_r(X) = 1$ at all other local nodes
- Use formulas for Lagrange interpolating polynomials on the el em en t

Integration over a reference element; element matrix

P1 elements and f(x) = x(1-x).

$$\begin{split} \tilde{A}_{0,0}^{(e)} &= \int_{-1}^{1} \tilde{\varphi}_{0}(X) \tilde{\varphi}_{0}(X) \frac{h}{2} dX \\ &= \int_{-1}^{1} \frac{1}{2} (1 - X) \frac{1}{2} (1 - X) \frac{h}{2} dX = \frac{h}{8} \int_{-1}^{1} (1 - X)^{2} dX = \frac{h}{3} \\ \tilde{A}_{1,0}^{(e)} &= \int_{-1}^{1} \tilde{\varphi}_{1}(X) \tilde{\varphi}_{0}(X) \frac{h}{2} dX \end{split}$$
(13

$$\begin{split} \widetilde{A}_{1,0}^{(e)} &= \int_{-1}^{1} \widetilde{\varphi}_{1}(X) \widetilde{\varphi}_{0}(X) \frac{h}{2} dX \\ &= \int_{-1}^{1} \frac{1}{2} (1+X) \frac{1}{2} (1-X) \frac{h}{2} dX = \frac{h}{8} \int_{-1}^{1} (1-X^{2}) dX = \frac{h}{6} \end{split}$$
(1)

$$\tilde{A}_{0,1}^{(e)} = \tilde{A}_{1,0}^{(e)} \tag{15}$$

$$\begin{split} \tilde{A}_{0,1}^{(e)} &= \tilde{A}_{1,0}^{(e)} \\ \tilde{A}_{1,1}^{(e)} &= \int_{-1}^{1} \tilde{\varphi}_1(X) \tilde{\varphi}_1(X) \frac{h}{2} dX \end{split}$$

Integration over a reference element; element vector

$$\tilde{b}_{0}^{(e)} = \int_{-1}^{1} f(x(X)) \tilde{\varphi}_{0}(X) \frac{h}{2} dX
= \int_{-1}^{1} (x_{m} + \frac{1}{2}hX) (1 - (x_{m} + \frac{1}{2}hX)) \frac{1}{2} (1 - X) \frac{h}{2} dX
= -\frac{1}{24}h^{3} + \frac{1}{6}h^{2}x_{m} - \frac{1}{12}h^{2} - \frac{1}{2}hx_{m}^{2} + \frac{1}{2}hx_{m}$$

$$\tilde{b}_{1}^{(e)} = \int_{-1}^{1} f(x(X)) \tilde{\varphi}_{1}(X) \frac{h}{2} dX
= \int_{-1}^{1} (x_{m} + \frac{1}{2}hX) (1 - (x_{m} + \frac{1}{2}hX)) \frac{1}{2} (1 + X) \frac{h}{2} dX
= -\frac{1}{24}h^{3} - \frac{1}{6}h^{2}x_{m} + \frac{1}{12}h^{2} - \frac{1}{2}hx_{m}^{2} + \frac{1}{2}hx_{m}$$
(18)

 x_m : element midpoint.

Implementation

- Coming functions appear in fe_approx1D.py
- Functions can operate in symbolic or numeric mode
- The code documents all steps in finite element calculations!

```
Compute the element matrix
```

```
def element_matrix(phi, Omega_e, symbolic=True):
    n = len(phi)
    A_e = sp.zeros((n, n))
    X = sp.Symbol('X')
    if symbolic:
        h = sp.Symbol('h')
    else:
        h = Omega_e[i] - Omega_e[0]
    detJ = h/2 * da/dI
    for r in range(n):
        for s in range(r, n):
            A_e[r,s] = sp.integrate(phi[r]*phi[s]*detJ, (X, -1, 1))
            A_e[s,r] = A_e[r,s]
```

Tedious calculations! Let's use symbolic software >>> import sympy as sp >>> x, x_m, h, X = sp.symbols('x x_m h X') >>> sp.integrate(h/8*(1-X)**2, (X, -1, 1)) h/3 >>> sp.integrate(h/8*(1+X)*(1-X), (X, -1, 1)) h/6 >>> x = x_m + h/2*X >>> b.0 = sp.integrate(h/4*x*(1-x)*(1-X), (X, -1, 1)) >>> print bolous print sp.latex(b_0, mode='plain') - \frac{1}{24} h^{23} + \frac{1}{24} h^{23} - \half h x_{1} \frac{1}{24} h^{23} + \half h x_{2} \frac{1}{24} h^{23} h^

def element_vector(f, phi, Omega_e, symbolic=True): n = len(phi) b_e = sp.zeros((n, 1)) # Make f a function of I I = sp.Symbol('X') if symbolic: h = sp.Symbol('x') else: h = Omega_e[i] - Omega_e[i] x = (Omega_e[i] + Omega_e[i])/2 + h/2*X # mapping f = f.subs('x', x) # substitute mapping formula for x detJ = h/2 # da/dI for r in range(n): b_e[i] = sp.integrate(f*phi[r]*detJ, (X, -1, 1)) return b_e Note f.subs('x', x): replace x by x(X) such that f contains X


```
>>> h, x = sp.symbols('h x')
>>> nodes = [0, h, 2*h]
>>> elements = [[0, 1], [1, 2]]
>>> phi = basis(a=1)
>>> f = x*(1-x)
>>> A

(h/3, h/6, 0]
(h/6, 2*h/3, h/6)
[ 0, h/6, h/3]
>>> b

[ h**2/6 - h***3/12]
[ h**2/7 - 17*h**3/6]
[ 5*h**2/6 - 17*h**3/12]
>>> c = A.LUsolve(b)
>>> c

[ 12*(7*h**2/12 - 35*h**3/72)/(7*h)]
[ 7*(4*h**2/7 - 23*h**3/21)/(2*h)]
```

The structure of the coefficient matrix

Note: do this by hand to understand what is going on!

General result: the coefficient matrix is sparse

- Sparse = most of the entries are zeros
- Below: P1 elements

Exemplifying the sparsity for P2 elements

$$A = \frac{h}{30} \begin{pmatrix} 4 & 2 & -1 & 0 & 0 & 0 & 0 & 0 & 0 \\ 2 & 16 & 2 & 0 & 0 & 0 & 0 & 0 & 0 & 0 \\ -1 & 2 & 8 & 2 & -1 & 0 & 0 & 0 & 0 & 0 \\ 0 & 0 & 2 & 16 & 2 & 0 & 0 & 0 & 0 & 0 \\ 0 & 0 & -1 & 2 & 8 & 2 & -1 & 0 & 0 & 0 \\ 0 & 0 & 0 & 0 & 2 & 16 & 2 & 0 & 0 & 0 \\ 0 & 0 & 0 & 0 & 0 & 2 & 16 & 2 & 0 & 0 \\ 0 & 0 & 0 & 0 & 0 & -1 & 2 & 8 & 2 & -1 \\ 0 & 0 & 0 & 0 & 0 & 0 & 0 & 2 & 16 & 2 \\ 0 & 0 & 0 & 0 & 0 & 0 & 0 & -1 & 2 & 4 \end{pmatrix}$$

Matrix sparsity pattern for regular/random numbering of P1 elements

- Left: number nodes and elements from left to right
- Right: number nodes and elements arbitrarily





Matrix sparsity pattern for regular/random numbering of P3 elements

- Left: number nodes and elements from left to right
- Right: number nodes and elements arbitrarily





Sparse matrix storage and solution

The minimum storage requirements for the coefficient matrix $A_{i,j}$:

- P1 elements: only 3 nonzero entries per row
- P2 elements: only 5 nonzero entries per row
- P3 elements: only 7 nonzero entries per row
- It is important to utilize sparse storage and sparse solvers
- In Python: scipy.sparse package

Approximate $f \sim x^9$ by various elements; code

Compute a mesh with N_e elements, basis functions of degree d, and approximate a given symbolic expression f(x) by a finite element expansion $u(x) = \sum_i c_i \varphi_i(x)$:

```
import sympy as sp
from fe_approxID import approximate
x = sp.Symbol('x')
approximate(f=x*(1-x)**8, symbolic=False, d=1, N_e=4)
approximate(f=x*(1-x)**8, symbolic=False, d=2, N_e=2)
approximate(f=x*(1-x)**8, symbolic=False, d=1, N_e=8)
approximate(f=x*(1-x)**8, symbolic=False, d=2, N_e=4)
```

Approximate $f \sim x^9$ by various elements; plot x^9 by various

Comparison of finite element and finite difference approximation

- Finite difference approximation u_i of a function f(x): simply choose $u_i = f(x_i)$
- This is the same as $u \approx \sum_i c_i \varphi_i$ + interpolation (see next slide)
- $u \approx \sum_i c_i \varphi_i + \mathsf{Galerkin/projection}$ or least squares method: must derive and solve a linear system
- ullet What is really the difference in the approximation u?

Interpolation/collocation with finite elements

Let $\{x_i\}_{i\in\mathcal{I}_s}$ be the nodes in the mesh. Collocation/interpolation means

$$u(x_i) = f(x_i), i \in \mathcal{I}_s,$$

which translates to

$$\sum_{j\in\mathcal{I}_s}c_j\varphi_j(x_i)=f(x_i),$$

but $\varphi_j(x_i)=0$ if $i\neq j$ so the sum collapses to one term $c_i\varphi_i(x_i)=c_i$, and we have the result

$$c_i = f(x_i)$$

Same result as the standard finite difference approach, but finite elements define u also between the x_i points

Galerkin/project and least squares vs collocation/interpolation or finite differences

- Scope: work with P1 elements
- Use projection/Galerkin or least squares (equivalent)
- Interpret the resulting linear system as finite difference equations

The P1 finite element machinery results in a linear system where equation no i is

$$\frac{h}{6}(u_{i-1} + 4u_i + u_{i+1}) = (f, \varphi_i)$$

Note:

- ullet We have used u_i for c_i to make notation similar to finite differences
- The finite difference counterpart is just $u_i = f_i$

Expressing the left-hand side in finite difference operator notation

Rewrite the left-hand side of finite element equation no i:

$$h(u_i + \frac{1}{6}(u_{i-1} - 2u_i + u_{i+1})) = [h(u + \frac{h^2}{6}D_x D_x u)]_i$$

This is the standard finite difference approximation of

$$h(u+\frac{h^2}{6}u'')$$

Treating the right-hand side; Trapezoidal rule

$$(f,\varphi_i) = \int_{x_{i-1}}^{x_i} f(x) \frac{1}{h} (x - x_{i-1}) dx + \int_{x_i}^{x_{i+1}} f(x) \frac{1}{h} (1 - (x - x_i)) dx$$

Cannot do much unless we specialize f or use *numerical integration*. Trapezoidal rule using the nodes:

$$(f,\varphi_i) = \int_{\Omega} f\varphi_i dx \approx h \frac{1}{2} (f(x_0)\varphi_i(x_0) + f(x_N)\varphi_i(x_N)) + h \sum_{j=1}^{N-1} f(x_j)\varphi_i(x_j)$$

 $\varphi_i(x_i) = \delta_{ii}$, so this formula collapses to one term:

$$(f,\varphi_i)\approx hf(x_i), \quad i=1,\ldots,N-1.$$

Same result as in collocation (interpolation) and the finite difference method!

Finite element approximation vs finite differences

With Trapezoidal integration of (f, φ_i) , the finite element method essentially solve

$$u + \frac{h^2}{6}u'' = f$$
, $u'(0) = u'(L) = 0$,

by the finite difference method

$$[u+\frac{h^2}{6}D_XD_Xu=f]_i$$

With Simpson integration of (f, φ_i) we essentially solve

$$[u+\frac{h^2}{6}D_x D_x u=\bar{f}]_i,$$

where

$$\bar{f}_i = \frac{1}{3}(f_{i-1/2} + f_i + f_{i+1/2})$$

Note: as $h \to 0$, $hu'' \to 0$ and $\bar{f}_i \to f_i$.

Limitations of the nodes and element concepts

So far,

- Nodes: points for defining φ_i and computing u values
- Elements: subdomain (containing a few nodes)
- This is a common notion of nodes and elements

One problem:

- Our algorithms need nodes at the element boundaries
- This is often not desirable, so we need to throw the nodes and elements arrays away and find a more generalized element concept

Treating the right-hand side; Simpson's rule

$$\int_{\Omega} g(x)dx \approx \frac{h}{6} \left(g(x_0) + 2 \sum_{j=1}^{N-1} g(x_j) + 4 \sum_{j=0}^{N-1} g(x_{j+\frac{1}{2}}) + f(x_{2N}) \right),$$

Our case: $g=f\varphi_i$. The sums collapse because $\varphi_i=0$ at most of the points.

$$(f,\varphi_i)pprox rac{h}{3}(f_{i-rac{1}{2}}+f_i+f_{i+rac{1}{2}})$$

Conclusions:

- While the finite difference method just samples f at x_i, the finite element method applies an average (smoothing) of f around x_i
- ullet On the left-hand side we have a term $\sim hu''$, and u'' also contribute to smoothing
- There is some inherent smoothing in the finite element method

Making finite elements behave as finite differences

- Can we adjust the finite element method so that we do not get the extra hu'' smoothing term and averaging of f?
- This allows finite elements to inherit (desired) properties of finite differences

Result:

- Compute all integrals by the Trapezoidal method and P1 elements
- Specifically, the coefficient matrix becomes diagonal ("lumped") - no linear system (!)
- ullet Loss of accuracy? The Trapezoidal rule has error $\mathcal{O}(h^2)$, the same as the approximation error in P1 elements

The generalized element concept has cells, vertices, nodes, and degrees of freedom

- We introduce *cell* for the subdomain that we up to now called element
- A cell has *vertices* (interval end points)
- Nodes are, almost as before, points where we want to compute unknown functions
- ullet Degrees of freedom is what the c_j represent (usually function values at nodes)

The concept of a finite element

- a reference cell in a local reference coordinate system
- $oldsymbol{Q}$ a set of basis functions $ilde{arphi}_r$ defined on the cell
- a set of degrees of freedom (e.g., function values) that uniquely determine the basis functions such that $\tilde{\varphi}_r=1$ for degree of freedom number r and $\tilde{\varphi}_r=0$ for all other degrees of freedom
- a mapping between local and global degree of freedom numbers (dof map)
- a geometric *mapping* of the reference cell onto to cell in the physical domain: $[-1,1] \Rightarrow [x_I,x_R]$

Basic data structures: vertices, cells, dof_map

- Cell vertex coordinates: vertices (equals nodes for P1 elements)
- Element vertices: cells[e][r] holds global vertex number of local vertex nor in element e (same as elements for P1 elements)
- dof_map[e,r] maps local dof r in element e to global dof number (same as elements for Pd elements)

The assembly process now applies dof_map:

```
A[dof_map[e][r], dof_map[e][s]] += A_e[r,s]
b[dof_map[e][r]] += b_e[r]
```


Example: P0 elements

Example: Same mesh, but u is piecewise constant in each cell (P0 element). Same vertices and cells, but

May think of one node in the middle of each element.

Note

We will hereafter work with cells, vertices, and dof_map.

Computing the error of the approximation; principles

$$L^2$$
 error: $||e||_{L^2} = \left(\int_{\Omega} e^2 dx\right)^{1/2}$

Accurate approximation of the integral:

- Sample u(x) at many points in each element (call u_glob, returns x and u)
- Use the Trapezoidal rule based on the samples
- It is important to integrate u accurately over the elements
- (In a finite difference method we would just sample the mesh point values)

Computing the error of the approximation; details

Not

We need a version of the Trapezoidal rule valid for non-uniformly spaced points:

$$\int_{\Omega} g(x) dx \approx \sum_{j=0}^{n-1} \frac{1}{2} (g(x_j) + g(x_{j+1})) (x_{j+1} - x_j)$$

How does the error depend on h and d?

Theory and experiments show that the least squares or projection/Galerkin method in combination with $P\emph{d}$ elements of equal length \emph{h} has an error

$$||e||_{L^2} = Ch^{d+1}$$

where C depends on f, but not on h or d.

Cubic Hermite polynomials; definition

• Can we construct $\varphi_i(x)$ with continuous derivatives? Yes!

Consider a reference cell [-1,1]. We introduce two nodes, X=-1 and X=1. The degrees of freedom are

- ullet 0: value of function at X=-1
- 1: value of first derivative at X=-1
- ullet 2: value of function at X=1
- 3: value of first derivative at X=1

Derivatives as unknowns ensure the same $\varphi_i'(x)$ value at nodes and thereby continuous derivatives.

Cubic Hermite polynomials; derivation

4 constraints on $\tilde{\varphi}_r$ (1 for dof r, 0 for all others):

•
$$\tilde{\varphi}_0(X_{(0)}) = 1$$
, $\tilde{\varphi}_0(X_{(1)}) = 0$, $\tilde{\varphi}'_0(X_{(0)}) = 0$, $\tilde{\varphi}'_0(X_{(1)}) = 0$

•
$$\tilde{\varphi}'_1(X_{(0)}) = 1$$
, $\tilde{\varphi}'_1(X_{(1)}) = 0$, $\tilde{\varphi}_1(X_{(0)}) = 0$, $\tilde{\varphi}_1(X_{(1)}) = 0$

$$\bullet \ \ \tilde{\varphi}_2(X_{(1)})=1, \ \tilde{\varphi}_2(X_{(0)})=0, \ \tilde{\varphi}_2'(X_{(0)})=0, \ \tilde{\varphi}_2'(X_{(1)})=0$$

•
$$\tilde{\varphi}_3'(X_{(1)}) = 1$$
, $\tilde{\varphi}_3'(X_{(0)}) = 0$, $\tilde{\varphi}_3(X_{(0)}) = 0$, $\tilde{\varphi}_3(X_{(1)}) = 0$

This gives 4 linear, coupled equations for each $\tilde{\varphi}_r$ to determine the 4 coefficients in the cubic polynomial

Cubic Hermite polynomials; result

$$\tilde{\varphi}_0(X) = 1 - \frac{3}{4}(X+1)^2 + \frac{1}{4}(X+1)^3$$
 (19)

$$\tilde{\varphi}_1(X) = -(X+1)(1-\frac{1}{2}(X+1))^2 \tag{20}$$

$$\tilde{\varphi}_2(X) = \frac{3}{4}(X+1)^2 - \frac{1}{2}(X+1)^3 \tag{21}$$

$$\tilde{\varphi}_3(X) = -\frac{1}{2}(X+1)(\frac{1}{2}(X+1)^2 - (X+1)) \tag{22}$$

(23)

Numerical integration

- $ullet \int_\Omega f arphi_i dx$ must in general be computed by numerical integration
- Numerical integration is often used for the matrix too

Common form of a numerical integration rule

$$\int_{-1}^{1} g(X)dX \approx \sum_{i=0}^{M} w_{i}g(\bar{X}_{i}),$$

wher

- \bar{X}_i are integration points
- w; are integration weights

Different rules correspond to different choices of points and weights

The Midpoint rule

Simplest possibility: the Midpoint rule,

$$\int_{-1}^{1} g(X)dX \approx 2g(0), \quad \bar{X}_{0} = 0, \ w_{0} = 2,$$

Exact for linear integrands

Newton-Cotes rules apply the nodes

- ullet ldea: use a fixed, uniformly distributed set of points in [-1,1]
- The points often coincides with nodes
- ullet Very useful for making $arphi_i arphi_j = 0$ and get diagonal ("mass") matrices ("lumping")

The Trapezoidal rule:

$$\int_{-1}^{1} g(X)dX \approx g(-1) + g(1), \quad \bar{X}_{0} = -1, \ \bar{X}_{1} = 1, \ w_{0} = w_{1} = 1,$$

Simpson's rule:

$$\int_{-1}^{1} g(X) dX \approx \frac{1}{3} (g(-1) + 4g(0) + g(1)),$$

wher

Gauss-Legendre rules apply optimized points

- Optimize the location of points to get higher accuracy
- Gauss-Legendre rules (quadrature) adjust points and weights to integrate polynomials exactly

$$M=1: \quad \bar{X}_0=-\frac{1}{\sqrt{3}}, \ \bar{X}_1=\frac{1}{\sqrt{3}}, \ w_0=w_1=1$$
 (24)

$$M=2:$$
 $\bar{X}_0=-\sqrt{\frac{3}{5}}, \ \bar{X}_0=0, \ \bar{X}_2=\sqrt{\frac{3}{5}}, \ w_0=w_2=\frac{5}{9}, \ w_1=\frac{8}{9}$ (25)

- M = 1: integrates 3rd degree polynomials exactly
- M=2: integrates 5th degree polynomials exactly
- In general, M-point rule integrates a polynomial of degree 2M+1 exactly.

See numint.py for a large collection of Gauss-Legendre rules.

Approximation of functions in 2D

Extensibility of 1D ideas.

All the concepts and algorithms developed for approximation of 1D functions f(x) can readily be extended to 2D functions f(x, y) and 3D functions f(x, y, z). Key formulas stay the same.

Quick overview of the 2D case

Inner product in 2D:

$$(f,g) = \int_{\Omega} f(x,y)g(x,y)dxdy$$

Least squares and project/Galerkin lead to a linear system

$$\sum_{j \in \mathcal{I}_s} A_{i,j} c_j = b_i, \quad i \in \mathcal{I}_s$$
$$A_{i,j} = (\psi_i, \psi_j)$$
$$b_i = (f, \psi_i)$$

Challenge: How to construct 2D basis functions $\psi_i(x, y)$?

2D basis functions as tensor products of 1D functions

Use a 1D basis for x variation and a similar for y variation:

$$V_x = \text{span}\{\hat{\psi}_0(x), \dots, \hat{\psi}_{N_x}(x)\}$$
 (26)

$$V_y = \text{span}\{\hat{\psi}_0(y), \dots, \hat{\psi}_{N_y}(y)\}$$
 (27)

The 2D vector space can be defined as a *tensor product* $V=V_x\otimes V_y$ with basis functions

$$\psi_{p,q}(x,y) = \hat{\psi}_p(x)\hat{\psi}_q(y) \quad p \in \mathcal{I}_x, q \in \mathcal{I}_y.$$

Tensor products

Given two vectors $a=(a_0,\ldots,a_M)$ and $b=(b_0,\ldots,b_N)$ their outer tensor product, also called the dyadic product, is $p=a\otimes b$, defined through

$$p_{i,j} = a_i b_j, \quad i = 0, \dots, M, \ j = 0, \dots, N.$$

Note: p has two indices (as a matrix or two-dimensional array)

Example: 2D basis as tensor product of 1D spaces,

$$\psi_{p,q}(x,y) = \hat{\psi}_p(x)\hat{\psi}_q(y), \quad p \in \mathcal{I}_x, q \in \mathcal{I}_y$$

Double or single index?

The 2D basis can employ a double index and double sum:

$$u = \sum_{p \in \mathcal{I}_x} \sum_{q \in \mathcal{I}_y} c_{p,q} \psi_{p,q}(x,y)$$

Or just a single index:

$$u = \sum_{j \in \mathcal{I}_s} c_j \psi_j(x, y)$$

with an index mapping $(p, q) \rightarrow i$:

$$\psi_i(x,y) = \hat{\psi}_p(x)\hat{\psi}_q(y), \quad i = pN_y + q \text{ or } i = qN_x + p$$

Example on 2D (bilinear) basis functions; formulas

In 1D we use the basis

$$\{1, x\}$$

2D tensor product (all combinations):

$$\psi_{0,0}=1, \quad \psi_{1,0}=x, \quad \psi_{0,1}=y, \quad \psi_{1,1}=xy$$

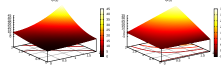
or with a single index:

$$\psi_0 = 1$$
, $\psi_1 = x$, $\psi_2 = y$, $\psi_3 = xy$

See notes for details of a hand-calculation.

Example on 2D (bilinear) basis functions; plot

Quadratic $f(x,y) = (1+x^2)(1+2y^2)$ (left), bilinear u (right):



Implementation; principal changes to the 1D code

Very small modification of approx1D.py:

- Omega = [[O, L_x], [O, L_y]]
- Symbolic integration in 2D
- Construction of 2D (tensor product) basis functions

Implementation; 2D basis functions

Tensor product of 1D "Taylor-style" polynomials x^i :

```
def taylor(x, y, Nx, Ny):
    return [x**i*y**j for i in range(Nx+1) for j in range(Ny+1)]
```

Tensor product of 1D sine functions $\sin((i+1)\pi x)$:

Complete code in approx2D.py

Implementation; application

$$f(x,y) = (1+x^2)(1+2y^2)$$
>>> from approx2D import *
>>> f = (1*x**2)*(1*2*y**2)
>>> psi = taylor(x, y, 1, 1)
>>> Omega = [[0, 2], [0, 2]]
>>> y, c = least_squares(f, psi, Omega)
>>> print u
8*x*y - 2*x/3 + 4*y/3 - 1/9
>>> print sp.expand(f)
2*x**2*y**2 + x**2 + 2*y**2 + 1

Implementation; trying a perfect expansion

Add higher powers to the basis such that $f \in V$:

Expected: u=f when $f\in V$

Generalization to 3D

Key idea:

$$V = V_x \otimes V_y \otimes V_z$$

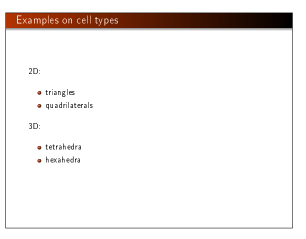
Repeated outer tensor product of multiple vectors

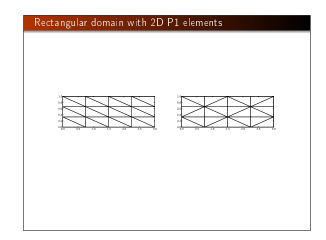
$$\begin{aligned} a^{(q)} &= (a_0^{(q)}, \dots, a_{N_q}^{(q)}), \quad q = 0, \dots, m \\ p &= a^{(0)} \otimes \dots \otimes a^{(m)} \\ p_{b_i, i_1, \dots, i_m} &= a_{i_1}^{(0)} a_{i_1}^{(1)} \dots a_{i_m}^{(m)} \end{aligned}$$

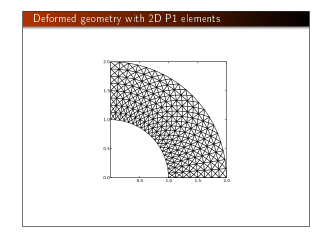
$$\psi_{p,q,r}(x,y,z) = \hat{\psi}_{p}(x)\hat{\psi}_{q}(y)\hat{\psi}_{r}(z)$$

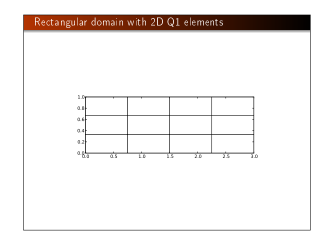
$$u(x,y,z) = \sum_{p \in \mathcal{I}_{x}} \sum_{q \in \mathcal{I}_{y}} \sum_{r \in \mathcal{I}_{z}} c_{p,q,r} \psi_{p,q,r}(x,y,z)$$

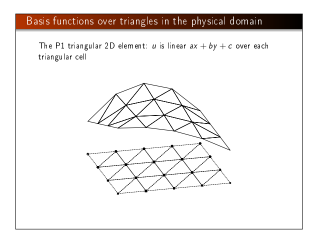
Finite elements in 2D and 3D The two great advantages of the finite element method: • Can handle complex-shaped domains in 2D and 3D • Can easily provide higher-order polynomials in the approximation Finite elements in 1D: mostly for learning, insight, debugging





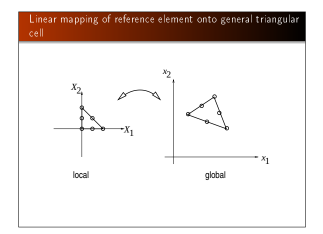






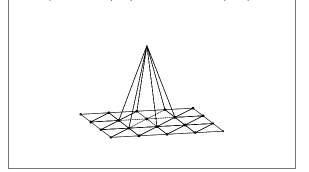
Basic features of 2D elements

- Cells = triangles
- Vertices = corners of the cells
- Nodes = vertices
- Degrees of freedom = function values at the nodes



φ_i : pyramid shape, composed of planes

- $\varphi_i(x,y)$ varies linearly over each cell
- $ullet arphi_i = 1$ at vertex (node) i, 0 at all other vertices (nodes)

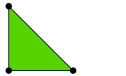


Element matrices and vectors

 As in 1D, the contribution from one cell to the matrix involves just a few entries, collected in the element matrix and vector

- $\varphi_i \varphi_j \neq 0$ only if i and j are degrees of freedom (vertices/nodes) in the same element
- ullet The 2D P1 element has a 3 imes 3 element matrix





$$\tilde{\varphi}_0(X,Y) = 1 - X - Y \tag{28}$$

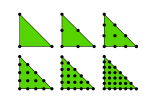
$$\tilde{\varphi}_1(X,Y) = X \tag{29}$$

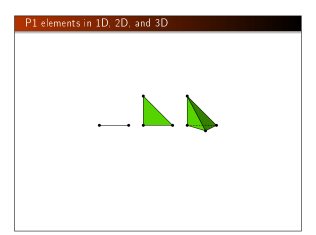
(30)

$$\tilde{\varphi}_2(X, Y) = Y$$

Higher-degree $ilde{arphi}_r$ introduce more nodes (dof = node values)

2D P1, P2, P3, P4, P5, and P6 elements





P2 elements in 1D, 2D, and 3D



- Interval, triangle, tetrahedron: simplex element (plural quick-form: simplices)
- Side of the cell is called face
- Thetrahedron has also edges

Affine mapping of the reference cell; formula

Mapping of local X = (X, Y) coordinates in the reference cell to global, physical x = (x, y) coordinates:

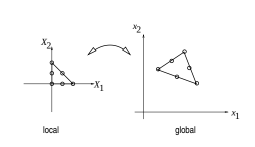
$$\mathbf{x} = \sum_{r} \tilde{\varphi}_r^{(1)}(\mathbf{X}) \mathbf{x}_{q(e,r)} \tag{31}$$

where

- r runs over the local vertex numbers in the cell
- x_i are the (x, y) coordinates of vertex i
- $\tilde{\varphi}_r^{(1)}$ are P1 basis functions

This mapping preserves the straight/planar faces and edges.

Affine mapping of the reference cell; figure

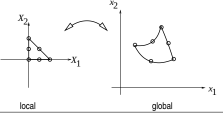


Isoparametric mapping of the reference cell

Idea: Use the basis functions of the element (not only the P1 functions) to map the element

$$x = \sum_{r} \tilde{\varphi}_r(X) x_{q(e,r)}$$

Advantage: higher-order polynomial basis functions now map the reference cell to a curved triangle or tetrahedron.



Computing integrals

Integrals must be transformed from $\Omega^{(e)}$ (physical cell) to $\tilde{\Omega}^r$ (reference cell):

$$\int_{\Omega^{(e)}} \varphi_i(\mathbf{x}) \varphi_j(\mathbf{x}) \, d\mathbf{x} = \int_{\tilde{\Omega}'} \tilde{\varphi}_i(\mathbf{X}) \tilde{\varphi}_j(\mathbf{X}) \, \det J \, d\mathbf{X}$$

$$\int_{\Omega^{(e)}} \varphi_i(\mathbf{x}) f(\mathbf{x}) \, d\mathbf{x} = \int_{\tilde{\Omega}'} \tilde{\varphi}_i(\mathbf{X}) f(\mathbf{x}(\mathbf{X})) \, \det J \, d\mathbf{X}$$
(32)

$$\int_{\Omega^{(e)}} \varphi_i(\mathbf{x}) f(\mathbf{x}) \, \mathrm{d}\mathbf{x} = \int_{\tilde{\Omega}'} \tilde{\varphi}_i(\mathbf{X}) f(\mathbf{x}(\mathbf{X})) \, \mathrm{det} \, J \, \mathrm{d}\mathbf{X}$$
 (33)

where dx = dxdy or dx = dxdydz and $\det J$ is the determinant of the Jacobian of the mapping x(X).

$$J = \left[\begin{array}{cc} \frac{\partial x}{\partial X} & \frac{\partial x}{\partial Y} \\ \frac{\partial y}{\partial X} & \frac{\partial y}{\partial Y} \end{array} \right], \quad \det J = \frac{\partial x}{\partial X} \frac{\partial y}{\partial Y} - \frac{\partial x}{\partial Y} \frac{\partial y}{\partial X}$$

Affine mapping (31): det $J = 2\Delta$, $\Delta = \text{cell volume}$

Remark on going from 1D to 2D/3D

Finite elements in 2D and 3D builds on the same ideas and concepts as in 1D, but there is simply much more to compute because the specific mathematical formulas in 2D and 3D are more complicated and the book keeping with dof maps also gets more complicated. The manual work is tedious, lengthy, and error-prone so automation by the computer is a must.

We shall apply least squares, Galerkin/projection, and collocation to differential equation models

Our aim is to extend the ideas for approximating f by u, or solving

$$u = f$$

to real, spatial differential equations like

$$-u'' + bu = f$$
, $u(0) = C$, $u'(L) = D$

Emphasis will be on the Galerkin/projection method

Abstract differential equation

$$\mathcal{L}(u) = 0, \quad x \in \Omega$$

Examples (1D problems):

$$\mathcal{L}(u) = \frac{d^2 u}{dx^2} - f(x),$$

$$\mathcal{L}(u) = \frac{d}{dx} \left(\alpha(x) \frac{du}{dx} \right) + f(x),$$

$$\mathcal{L}(u) = \frac{d}{dx} \left(\alpha(u) \frac{du}{dx} \right) - au + f(x),$$

$$\mathcal{L}(u) = \frac{d}{dx} \left(\alpha(u) \frac{du}{dx} \right) + f(u, x)$$

Abstract boundary conditions

$$\mathcal{B}_0(u) = 0, \ x = 0, \quad \mathcal{B}_1(u) = 0, \ x = L$$

Examples:

$$B_i(u) = u - g$$

$$\mathcal{B}_i(u) = -\alpha \frac{du}{dx} - g$$

$$\mathcal{B}_i(u) = u - g,$$
 Dirichlet condition $\mathcal{B}_i(u) = -lpha rac{du}{dx} - g,$ Neumann condition $\mathcal{B}_i(u) = -lpha rac{du}{dx} - h(u - g),$ Robin condition

Reminder about notation

- $u_e(x)$ is the symbol for the exact solution of $\mathcal{L}(u_e) = 0$ + $B_i = 0$
- u(x) denotes an approximate solution
- $V = \operatorname{span}\{\psi_0(x), \dots, \psi_N(x)\}$, V has basis $\{\psi_i\}_{i \in \mathcal{I}_s}$
- We seek $u \in V$
- $\mathcal{I}_s = \{0, \dots, N\}$ is an index set
- $u(x) = \sum_{j \in \mathcal{I}_s} c_j \psi_j(x)$
- Inner product: $(u, v) = \int_{\Omega} uv \, dx$
- Norm: $||u|| = \sqrt{(u,u)}$

New topics: variational formulation and boundary conditions

Much is similar to approximating a function (solving u = f), but two new topics are needed:

- Variational formulation of the differential equation problem (including integration by parts)
- Handling of boundary conditions

Residual-minimizing principles

- ullet When solving u=f we knew the error e=f-u and could use principles for minimizing the error
- ullet When solving $\mathcal{L}(u_{\mathrm{e}})=0$ we do not know u_{e} and cannot work with the error $e=u_{\mathrm{e}}-u$
- ullet We can only know the error in the equation: the residual R

Inserting $u=\sum_{i}c_{j}\psi_{j}$ in $\mathcal{L}=0$ gives a residual R

$$\mathcal{L}(u) = \mathcal{L}(\sum_{j} c_{j}\psi_{j}) = R \neq 0$$

Goal: minimize R with respect to $\{c_i\}_{i\in\mathcal{I}_{\mathbf{z}}}$ (and hope it makes a small e too)

$$R = R(c_0, \ldots, c_N; x)$$

The least squares method

ldea: minimize

$$E = ||R||^2 = (R, R) = \int_{\Omega} R^2 dx$$

Minimization wrt $\left\{c_i
ight\}_{i\in\mathcal{I}_s}$ implies

$$\frac{\partial E}{\partial c_i} = \int_{\Omega} 2R \frac{\partial R}{\partial c_i} dx = 0 \quad \Leftrightarrow \quad (R, \frac{\partial R}{\partial c_i}) = 0, \quad i \in \mathcal{I}_s$$

 $\mathit{N}+1$ equations for $\mathit{N}+1$ unknowns $\{c_i\}_{i\in\mathcal{I}_r}$

The Galerkin method

Idea: make R orthogonal to V,

$$(R, v) = 0, \quad \forall v \in V$$

This implies

$$(R, \psi_i) = 0, \quad i \in \mathcal{I}_s$$

N+1 equations for N+1 unknowns $\{c_i\}_{i\in\mathcal{I}_e}$

The Method of Weighted Residuals

Generalization of the Galerkin method: demand R orthogonal to some space W, possibly $W \neq V$:

$$(R, v) = 0, \forall v \in W$$

If $\{w_0, \ldots, w_N\}$ is a basis for W:

$$(R, w_i) = 0, \quad i \in \mathcal{I}_s$$

- ullet N+1 equations for N+1 unknowns $\{c_i\}_{i\in\mathcal{I}_{ullet}}$
- Weighted residual with $w_i = \partial R/\partial c_i$ gives least squares

New terminology: test and trial functions

- ullet ψ_j used in $\sum_i c_j \psi_j$ is called *trial function*
- ullet ψ_i or w_i used as weight in Galerkin's method is called test

The collocation method

Idea: demand R=0 at N+1 points in space

$$R(x_i; c_0, \ldots, c_N) = 0, \quad i \in \mathcal{I}_s$$

The collocation method is a weighted residual method with delta functions as weights

$$0 = \int_{\Omega} R(x; c_0, \ldots, c_N) \delta(x - x_i) dx = R(x_i; c_0, \ldots, c_N)$$

property of
$$\delta(x)$$
: $\int_{\Omega} f(x)\delta(x-x_i)dx = f(x_i), \quad x_i \in \Omega$



Examples on using the principles

Goal

Exemplify the least squares, Galerkin, and collocation methods in a simple 1D problem with global basis functions.

The first model problem

$$-u''(x) = f(x), \quad x \in \Omega = [0, L], \quad u(0) = 0, \ u(L) = 0$$

Basis functions:

$$\psi_i(x) = \sin\left((i+1)\pi\frac{x}{L}\right), \quad i \in \mathcal{I}_s$$

Residual:

$$R(x; c_0, ..., c_N) = u''(x) + f(x),$$

$$= \frac{d^2}{dx^2} \left(\sum_{j \in \mathcal{I}_s} c_j \psi_j(x) \right) + f(x),$$

$$= -\sum_{j \in \mathcal{I}_s} c_j \psi_j''(x) + f(x)$$

Boundary conditions

Since u(0)=u(L)=0 we must ensure that all $\psi_i(0)=\psi_i(L)=0$, because then

$$u(0) = \sum_{j} c_{j} \psi_{j}(0) = 0, \quad u(L) = \sum_{j} c_{j} \psi_{j}(L) = 0$$

- u known: Dirichlet boundary condition
- u' known: Neumann boundary condition
- Must have $\psi_i = 0$ where Dirichlet conditions apply

The least squares method; principle

$$(R, \frac{\partial R}{\partial c_i}) = 0, \quad i \in \mathcal{I}_s$$

$$\frac{\partial R}{\partial c_i} = \frac{\partial}{\partial c_i} \left(\sum_{j \in \mathcal{I}_s} c_j \psi_j''(x) + f(x) \right) = \psi_i''(x)$$

Because

$$\frac{\partial}{\partial c_i} \left(c_0 \psi_0'' + c_1 \psi_1'' + \dots + c_{i-1} \psi_{i-1}'' + \frac{c_i \psi_i''}{i} + c_{i+1} \psi_{i+1}'' + \dots + c_N \psi_N'' \right) =$$

The least squares method; equation system

$$(\sum_{j} c_{j}\psi_{j}'' + f, \psi_{i}'') = 0, \quad i \in \mathcal{I}_{s}$$

Rearrangement:

$$\sum_{j\in\mathcal{I}_s} (\psi_i'', \psi_j'') c_j = -(f, \psi_i''), \quad i\in\mathcal{I}_s$$

This is a linear system

$$\sum_{i\in\mathcal{I}_s}A_{i,j}c_j=b_i,\quad i\in\mathcal{I}_s$$

The least squares method; matrix and right-hand side expressions

$$\begin{split} A_{i,j} &= (\psi_i'', \psi_j'') \\ &= \pi^4 (i+1)^2 (j+1)^2 L^{-4} \int_0^L \sin\left((i+1)\pi\frac{x}{L}\right) \sin\left((j+1)\pi\frac{x}{L}\right) \, dx \\ &= \left\{ \begin{array}{l} \frac{1}{2} L^{-3} \pi^4 (i+1)^4 & i = j \\ 0, & i \neq j \end{array} \right. \\ b_i &= -(f, \psi_i'') = (i+1)^2 \pi^2 L^{-2} \int_0^L f(x) \sin\left((i+1)\pi\frac{x}{L}\right) \, dx \end{split}$$

Orthogonality of the basis functions gives diagonal matrix

Useful property of the chosen basis functions:

$$\int\limits_{0}^{L} \sin\left((i+1)\pi\frac{\mathbf{x}}{L}\right) \sin\left((j+1)\pi\frac{\mathbf{x}}{L}\right) \, d\mathbf{x} = \delta_{ij}, \qquad \delta_{ij} = \left\{ \begin{array}{l} \frac{1}{2}L & i=j\\ 0, & i\neq j \end{array} \right.$$

 \Rightarrow $(\psi_i'', \psi_i'') = \delta_{ij}$, i.e., diagonal $A_{i,j}$, and we can easily solve for c_i :

$$c_i = \frac{2L}{\pi^2(i+1)^2} \int_0^L f(x) \sin\left((i+1)\pi\frac{x}{L}\right) dx$$

Least squares method; solution

Let sympy do the work (f(x) = 2):

$$c_i = 4\frac{L^2\left(\left(-1\right)^i + 1\right)}{\pi^3\left(i^3 + 3i^2 + 3i + 1\right)}, \quad u(x) = \sum_{k=0}^{N/2} \frac{8L^2}{\pi^3(2k+1)^3} \sin\left((2k+1)\pi\frac{x}{L}\right)$$

Fast decay: $c_2=c_0/27$, $c_4=c_0/125$ - only one term might be good enough:

$$u(x) \approx \frac{8L^2}{3} \sin\left(\pi \frac{x}{L}\right)$$

The Galerkin method; principle

$$R = u'' + f$$
:

$$(u'' + f, v) = 0, \forall v \in V,$$

or rearranged,

$$(u'', v) = -(f, v), \forall v \in V$$

This is a variational formulation of the differential equation problem.

 $\forall v \in V$ is equivalent with $\forall v \in \psi_i, i \in \mathcal{I}_s$:

$$(\sum_{i \in \mathcal{I}} c_j \psi_j'', \psi_i) = -(f, \psi_i), \quad i \in \mathcal{I}_s$$

The Galerkin method; solution

Since $\psi_i'' \propto \psi_i$, Galerkin's method gives the same linear system and the same solution as the least squares method (in this particular example).

The collocation method

 ${\it R}=0$ (i.e.,the differential equation) must be satisfied at ${\it N}+1$ points:

$$-\sum_{j\in\mathcal{I}_s}c_j\psi_j''(x_i)=f(x_i),\quad i\in\mathcal{I}_s$$

This is a linear system $\sum_{i} A_{i,j} = b_i$ with entries

$$A_{i,j} = -\psi_j''(x_i) = (j+1)^2 \pi^2 L^{-2} \sin\left((j+1)\pi \frac{x_i}{L}\right), \quad b_i = 2$$

Choose: N = 0, $x_0 = L/2$

$$c_0=2L^2/\pi^2$$

Comparison of the methods

- Exact solution: u(x) = x(L-x)
- Galerkin or least squares (N=0): $u(x)=8L^2\pi^{-3}\sin(\pi x/L)$
- Collocation method (N=0): $u(x)=2L^2\pi^{-2}\sin(\pi x/L)$.

```
>>> import sympy as sp
>>> f Computing with Dirichlet conditions: -u''-2 and sines
>>> x, L = sp. symbols('x L')
>>> e.Galerkin = x*(L-x) - 3*L**2*sp.pi**(-3)*sp.sin(sp.pi*x/L)
>>> e.colloc = x*(L-x) - 2*L**2*sp.pi**(-2)*sp.sin(sp.pi*x/L)
>>> f Verify max error for x=L/2
>>> dedx_Galerkin = sp.diff(e.Galerkin, x)
>>> dedx_Galerkin.subs(x, L/2)
0
>>> dedx_colloc = sp.diff(e.colloc, x)
>>> dedx_colloc subs(x, L/2)
0

# Compute max error: x=L/2, evaluate numerical, and simplify
-0.00812*el**2
>>> sp.simplify(e_Galerkin.subs(x, L/2).evalf(n=3))
-0.00812*el**2
>>> sp.simplify(e_colloc.subs(x, L/2).evalf(n=3))
0.0473*L**2
```

Integration by parts has many advantages

Second-order derivatives will hereafter be integrated by parts

$$\int_0^L u''(x)v(x)dx = -\int_0^L u'(x)v'(x)dx + [vu']_0^L$$
$$= -\int_0^L u'(x)v'(x)dx + u'(L)v(L) - u'(0)v(0)$$

Motivation:

- Lowers the order of derivatives
- Gives more symmetric forms (incl. matrices)
- Enables easy handling of Neumann boundary conditions
- \bullet Finite element basis functions φ_i have discontinuous derivatives (at cell boundaries) and are not suited for terms with φ_i''

We use a boundary function to deal with non-zero Dirichlet boundary conditions

- What about nonzero Dirichlet conditions? Say u(L) = D
- We always require $\psi_i(L) = 0$ (i.e., $\psi_i = 0$ where Dirichlet conditions applies)
- ullet Problem: $u(L) = \sum_j c_j \psi_j(L) = \sum_j c_j \cdot 0 = 0
 eq D$ always!
- Solution: $u(x) = B(x) + \sum_{i} c_{i} \psi_{i}(x)$
- ullet B(x): user-constructed boundary function that fulfills the Dirichlet conditions
- If u(L) = D, make sure B(L) = D
- ullet No restrictions of how B(x) varies in the interior of Ω

Example on constructing a boundary function for two Dirichlet conditions

Dirichlet conditions: u(0) = C and u(L) = D. Choose for example

$$B(x) = \frac{1}{L}(C(L-x) + Dx):$$
 $B(0) = C, B(L) = D$

$$u(x) = B(x) + \sum_{j \in \mathcal{I}_s} c_j \psi_j(x),$$

$$u(0) = B(0) = C, \quad u(L) = B(L) = D$$

Example on constructing a boundary function for one Dirichlet conditions

Dirichlet condition: u(L) = D. Choose for example

$$B(x) = D$$
: $B(L) = D$

$$u(x) = B(x) + \sum_{j \in \mathcal{I}_s} c_j \psi_j(x),$$

$$u(L) = B(L) = D$$

With a B(x), $u \notin V$, but $\sum_i c_i \psi_i \in V$

- ullet $\{\psi_i\}_{i\in\mathcal{I}_{\mathbf{s}}}$ is a basis for V
- $\sum_{j\in\mathcal{I}_s} c_j \psi_j(x) \in V$
- But $u \notin V!$
- Reason: say u(0) = C and $u \in V$ (any $v \in V$ has v(0) = C, then $2u \notin V$ because 2u(0) = 2C
- $\begin{array}{l} \bullet \text{ When } u(x) = B(x) + \sum_{j \in \mathcal{I}_s} c_j \psi_j(x), \ B \neq \emptyset, \ B \not \in V \text{ (in general) and } u \not \in V, \text{ but } (u-B) \in V \text{ since } \sum_j c_j \psi_j \in V \end{array}$

Abstract notation for variational formulations

The finite element literature (and much FEniCS documentation) applies an abstract notation for the variational formulation:

Find
$$(u - B) \in V$$
 such that

$$a(u, v) = L(v) \quad \forall v \in V$$

Example on abstract notation

$$-u'' = f$$
, $u'(0) = C$, $u(L) = D$, $u = D + \sum_{j} c_{j}\psi_{j}$

Variational formulation:

$$\int_{\Omega} u'v'dx = \int_{\Omega} fvdx - v(0)C \quad \text{or} \quad (u',v') = (f,v) - v(0)C \quad \forall v \in V$$

Abstract formulation: find $(u-B) \in V$ such that

$$a(u, v) = L(v) \quad \forall v \in V$$

We identify

$$a(u, v) = (u', v'), L(v) = (f, v) - v(0)C$$

Bilinear and linear forms

- a(u, v) is a bilinear form
- L(v) is a linear form

Linear form means

$$L(\alpha_1 v_1 + \alpha_2 v_2) = \alpha_1 L(v_1) + \alpha_2 L(v_2),$$

Bilinear form means

$$a(\alpha_1 u_1 + \alpha_2 u_2, v) = \alpha_1 a(u_1, v) + \alpha_2 a(u_2, v),$$

 $a(u, \alpha_1 v_1 + \alpha_2 v_2) = \alpha_1 a(u, v_1) + \alpha_2 a(u, v_2)$

In nonlinear problems: Find $(u-B) \in V$ such that $F(u;v)=0 \ \forall v \in V$

The linear system associated with abstract form

$$a(u, v) = L(v) \quad \forall v \in V \quad \Leftrightarrow \quad a(u, \psi_i) = L(\psi_i) \quad i \in \mathcal{I}_s$$

We can now derive the corresponding linear system once and for all by inserting $u=B+\sum_i c_i\psi_i$:

$$a(B + \sum_{j \in \mathcal{I}_s} c_j \psi_j, \psi_i) c_j = L(\psi_i) \quad i \in \mathcal{I}_s$$

Because of linearity,

$$\sum_{j \in \mathcal{I}_{\mathtt{s}}} \underbrace{a(\psi_{j}, \psi_{i})}_{A_{i,i}} c_{j} = \underbrace{L(\psi_{i}) - a(B, \psi_{i})}_{b_{i}} \quad i \in \mathcal{I}_{\mathtt{s}}$$

Equivalence with minimization problem

If a is symmetric: a(u, v) = a(v, u),

$$a(u, v) = L(v) \quad \forall v \in V$$

is equivalent to minimizing the functional

$$F(v) = \frac{1}{2}a(v,v) - L(v)$$

over all functions $v \in V$. That is,

$$F(u) \le F(v) \quad \forall v \in V$$

- Much used in the early days of finite elements
- Still much used in structural analysis and elasticity
- Not as general as Galerkin's method (since we require a(u, v) = a(v, u))

Examples on variational formulations

Goa

Derive variational formulations for many prototype differential equations in 1D that include

- variable coefficients
- mixed Dirichlet and Neumann conditions
- nonlinear coefficients

Variable coefficient; problem

$$-\frac{d}{dx}\left(\alpha(x)\frac{du}{dx}\right) = f(x), \quad x \in \Omega = [0, L], \ u(0) = C, \ u(L) = D$$

- Variable coefficient $\alpha(x)$
- Nonzero Dirichlet conditions at x = 0 and x = L
- Must have $\psi_i(0) = \psi_i(L) = 0$
- $V = \operatorname{span}\{\psi_0, \dots, \psi_N\}$
- $v \in V$: v(0) = v(L) = 0

$$u(x) = B(x) + \sum_{j \in \mathcal{I}_s} c_j \psi_j(x), \quad B(x) = C + \frac{1}{L}(D - C)x$$

Variable coefficient; Galerkin principle

$$R = -\frac{d}{dx} \left(a \frac{du}{dx} \right) - f$$

Galerkin's method:

$$(R, v) = 0, \forall v \in V$$

or with integrals:

$$\int_{\Omega} \left(\frac{d}{dx} \left(\alpha \frac{du}{dx} \right) - f \right) v \, dx = 0, \quad \forall v \in V$$

Variable coefficient; integration by parts

$$-\int_{\Omega} \frac{d}{dx} \left(\alpha(x) \frac{du}{dx} \right) v \, dx = \int_{\Omega} \alpha(x) \frac{du}{dx} \frac{dv}{dx} \, dx - \left[\alpha \frac{du}{dx} v \right]_{0}^{L}$$

Boundary terms vanish since v(0) = v(L) = 0

Variable coefficient; variational formulation

Variational formulation

Find $(u - B) \in V$ such that

$$\int_{\Omega} \alpha(x) \frac{du}{dx} \frac{dv}{dx} dx = \int_{\Omega} f(x) v dx, \quad \forall v \in V$$

Compact notation:

$$\underbrace{(\alpha u', v')}_{a(u,v)} = \underbrace{(f,v)}_{L(v)}, \quad \forall v \in V$$

Variable coefficient; linear system (the easy way)

With

$$a(u, v) = (\alpha u', v'), L(v) = (f, v)$$

we can just use the formula for the linear system:

$$\begin{aligned} A_{i,j} &= \mathsf{a}(\psi_j, \psi_i) = (\alpha \psi_j', \psi_i') = \int_{\Omega} \alpha \psi_j' \psi_i' \, \mathrm{d}x = \int_{\Omega} \psi_i' \alpha \psi_j' \, \mathrm{d}x \quad (= \mathsf{a}(\psi_i, \psi_j)) \\ b_i &= (f, \psi_i) - (\alpha B', \psi_i) = \int_{\Omega} (f \psi_i - \alpha L^{-1} (D - C) \psi_i') \, \mathrm{d}x \end{aligned}$$

Variable coefficient; linear system (full derivation)

 $v=\psi_i$ and $u=B+\sum_j c_j\psi_j$:

$$(\alpha B' + \alpha \sum_{j \in \mathcal{I}_s} c_j \psi'_j, \psi'_i) = (f, \psi_i), \quad i \in \mathcal{I}_s$$

Reorder to form linear system:

$$\sum_{j\in\mathcal{I}_{\mathtt{S}}}(\alpha\psi_{j}',\psi_{i}')c_{j}=(f,\psi_{i})+(\mathsf{a}\mathsf{L}^{-1}(\mathsf{D}-\mathsf{C}),\psi_{i}'),\quad i\in\mathcal{I}_{\mathtt{S}}$$

This is $\sum_{i} A_{i,j} c_j = b_i$ with

$$A_{i,j} = (a\psi'_j, \psi'_i) = \int_{\Omega} \alpha(x)\psi'_j(x)\psi'_i(x) dx$$

$$b_i = (f, \psi_i) + (aL^{-1}(D - C), \psi'_i) = \int_{\Omega} \left(f(x)\psi_i(x) + \alpha(x) \frac{D - C}{L} \psi'_i(x) \right)$$

First-order derivative in the equation and boundary condition; problem

$$-u''(x)+bu'(x)=f(x), \quad x\in\Omega=[0,L],\ u(0)=C,\ u'(L)=E$$

New features:

- ullet first-order derivative u' in the equation
- boundary condition with u': u'(L) = E

Initial steps:

- Must force $\psi_i(0) = 0$ because of Dirichlet condition at x = 0
- Boundary function: B(x) = C(L-x) or just B(x) = C
- ullet No requirements on $\psi_i(L)$ (no Dirichlet condition at x=L)

First-order derivative in the equation and boundary condition; details

$$u = C + \sum_{j \in \mathcal{I}_s} c_j \psi_i(x)$$

Galerkin's method: multiply by $\nu_{\rm t}$ integrate over $\Omega_{\rm t}$ integrate by parts.

$$(-u'' + bu' - f, v) = 0, \quad \forall v \in V$$

$$(u', v') + (bu', v) = (f, v) + [u'v]_0^L, \quad \forall v \in V$$

Now,
$$[u'v]_0^L = u'(L)v(L) = Ev(L)$$
 because $v(0) = 0$ and $u'(L) = E$:

$$(u'v')+(bu',v)=(f,v)+Ev(L), \forall v \in V$$

First-order derivative in the equation and boundary condition; observations

$$(u'v') + (bu', v) = (f, v) + Ev(L), \quad \forall v \in V,$$

Important:

- The boundary term can be used to implement Neumann conditions
- Forgetting the boundary term implies the condition u'=0 (!)
- Such conditions are called natural boundary conditions

First-order derivative in the equation and boundary condition; abstract notation

Abstract notation:

$$a(u, v) = L(v) \quad \forall v \in V$$

Here:

$$a(u, v) = (u', v') + (bu', v)$$

 $L(v) = (f, v) + Ev(L)$

First-order derivative in the equation and boundary condition; linear system

Insert $u = C + \sum_{j} c_{j} \psi_{j}$ and $v = \psi_{i}$:

$$\sum_{j\in\mathcal{I}_s} \underbrace{((\psi_j',\psi_i') + (b\psi_j',\psi_i))}_{A_{i,i}} c_j = \underbrace{(f,\psi_i) + E\psi_i(L)}_{b_i}$$

Observation: $A_{i,j}$ is not symmetric because of the term

$$(b\psi_j',\psi_i)=\int_{\Omega}b\psi_j'\psi_idx
eq \int_{\Omega}b\psi_i'\psi_jdx=(\psi_i',b\psi_j)$$

Terminology: natural and essential boundary conditions

$$(u', v') + (bu', v) = (f, v) + u'(L)v(L) - u'(0)v(0)$$

- Note: forgetting the boundary terms implies u'(L) = u'(0) = 0 (unless prescribe a Dirichlet condition)
- Conditions on *u'* are simply inserted in the variational form and called *natural conditions*
- ullet Conditions on u at x=0 requires modifying V (through $\psi_i(0)=0$) and are known as essential conditions

Lesson learned

It is easy to forget the boundary term when integrating by parts. That mistake may prescribe a condition on $u^\prime!$

Nonlinear coefficient; problem

Problem:

$$-(\alpha(u)u')' = f(u), \quad x \in [0, L], \ u(0) = 0, \ u'(L) = E$$

- V: basis $\{\psi_i\}_{i\in\mathcal{I}_s}$ with $\psi_i(0)=0$ because of u(0)=0
- How does the nonlinear coefficients $\alpha(u)$ and f(u) impact the variational formulation?
- (Not much!)

Nonlinear coefficient; variational formulation

Galerkin: multiply by v, integrate, integrate by parts

$$\int_0^L \alpha(u) \frac{du}{dx} \frac{dv}{dx} dx = \int_0^L f(u) v dx + [\alpha(u)vu']_0^L \quad \forall v \in V$$

- $\alpha(u(0))v(0)u'(0) = 0$ since v(0)
- $\alpha(u(L))v(L)u'(L) = \alpha(u(L))v(L)E$ since u'(L) = E

$$\int_0^L \alpha(u) \frac{du}{dx} \frac{dv}{dx} v \, dx = \int_0^L f(u) v \, dx + \alpha(u(L)) v(L) E \quad \forall v \in V$$

or

$$(\alpha(u)u',v')=(f(u),v)+\alpha(u(L))v(L)E \quad \forall v \in V$$

Nonlinear coefficient; where does the nonlinearity cause challenges?

- ullet Abstract notation: no a(u,v) and L(v) because a and L get nonlinear
- Instead: $F(u; v) = 0 \ \forall v \in V$
- What about forming a linear system? We get a nonlinear system of algebraic equations
- Must use methods like Picard iteration or Newton's method to solve nonlinear algebraic equations
- But: the variational formulation was not much affected by nonlinearities

Dirichlet and Neumann conditions; problem

$$-u''(x) = f(x), \quad x \in \Omega = [0,1], \quad u'(0) = C, \ u(1) = D$$

- Use a global polynomial basis $\psi_i \sim x^i$ on [0,1]
- Because of u(1) = D: $\psi_i(1) = 0$
- Basis: $\psi_i(x) = (1-x)^{i+1}, i \in \mathcal{I}_s$
- B(x) = Dx

Dirichlet and Neumann conditions; details (1)

$$A_{i,j} = (\psi'_j, \psi'_i) = \int_0^1 \psi'_i(x)\psi'_j(x)dx = \int_0^1 (i+1)(j+1)(1-x)^{i+j}dx,$$

Choose f(x) = 2:

$$b_i = (2, \psi_i) - (D, \psi_i') - C\psi_i(0)$$

= $\int_0^1 (2(1-x)^{i+1} - D(i+1)(1-x)^i) dx - C\psi_i(0)$

Dirichlet and Neumann conditions; details (2)

Can easily do the integrals with sympy. N=1:

$$\left(\begin{array}{cc} 1 & 1 \\ 1 & 4/3 \end{array}\right) \left(\begin{array}{c} c_0 \\ c_1 \end{array}\right) = \left(\begin{array}{c} -C + D + 1 \\ 2/3 - C + D \end{array}\right)$$

$$c_0 = -C + D + 2, \quad c_1 = -1,$$

$$u(x) = 1 - x^2 + D + C(x - 1) \quad (exact solution)$$

When the numerical method is exact

Assume that apart from boundary conditions, $u_{\rm e}$ lies in the same space V as where we seek u:

$$u = B + F, \quad F \in V$$
$$a(B + F, v) = L(v) \quad \forall v \in V$$
$$u_{e} = B + E, \quad E \in V$$

$$a(B+E,v)=L(v) \quad \forall v \in V$$

Subtract: $a(F - E, v) = 0 \implies E = F$ and $u = u_e$

Computing with finite elements

Tasks:

- Address the model problem -u''(x) = 2, u(0) = u(L) = 0
- Uniform finite element mesh with P1 elements
- Show all finite element computations in detail

Variational formulation

$$-u''(x) = 2$$
, $x \in (0, L)$, $u(0) = u(L) = 0$,

Variational formulation

$$(u', v') = (2, v) \quad \forall v \in V$$

How to deal with the boundary conditions?

Since u(0) = 0 and u(L) = 0, we must force

$$v(0) = v(L) = 0, \quad \psi_i(0) = \psi_i(L) = 0$$

Now we choose the finite element basis: $\psi_i = \varphi_i, i = 0, \dots, N_n$

Problem: $\varphi_0(0) \neq 0$ and $\varphi_{N_n}(L) \neq 0$

Solution: we just exclude φ_0 and φ_{N_0} from the basis and work with

$$\psi_i = \varphi_{i+1}, \quad i = 0, \dots, N = N_n - 2$$

Introduce index mapping u(j): $\psi_i = arphi_{
u(i)}$

$$u = \sum_{i \in \mathcal{I}_*} c_j \varphi_{\nu(i)}, \quad i = 0, \dots, N, \quad \nu(j) = j + 1$$

Irregular numbering: more complicated u(j) table

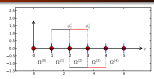
Computation in the global physical domain; formulas

$$A_{i,j} = \int_0^L \varphi'_{i+1}(x) \varphi'_{j+1}(x) dx, \quad b_i = \int_0^L 2\varphi_{i+1}(x) dx$$

Many will prefer to change indices to obtain a $\varphi_i'\varphi_j'$ product: $i+1 \to i, \ j+1 \to j$

$$A_{i-1,j-1} = \int_0^L \varphi_i'(x) \varphi_j'(x) dx, \quad b_{i-1} = \int_0^L 2\varphi_i(x) dx$$

Computation in the global physical domain; details



$$\varphi_i = \pm h^{-1}$$

$$A_{i-1,i-1} = h^{-2}2h = 2h^{-1}, \quad A_{i-1,i-2} = h^{-1}(-h^{-1})h = -h^{-1}, \quad A_{i-1,i} = -h^{-1}$$

$$b_{i-1} = 2(\frac{1}{2}h + \frac{1}{2}h) = 2h$$

Computation in the global physical domain; linear system

Write out the corresponding difference equation

General equation at node i:

$$-\frac{1}{h}c_{i-1} + \frac{2}{h}c_i - \frac{1}{h}c_{i+1} = 2h$$

Now, $c_i = u(x_{i+1}) \equiv u_{i+1}$. Writing out the equation at node i-1,

$$-\frac{1}{h}c_{i-2} + \frac{2}{h}c_{i-1} - \frac{1}{h}c_i = 2h$$

translates directly to

$$-\frac{1}{h}u_{i-1} + \frac{2}{h}u_i - \frac{1}{h}u_{i+1} = 2h$$

Comparison with a finite difference discretization

The standard finite difference method for -u''=2 is

$$-\frac{1}{h^2}u_{i-1} + \frac{2}{h^2}u_i - \frac{1}{h^2}u_{i+1} = 2$$

Multiply by h!

The finite element method and the finite difference method are identical *in this example*.

(Remains to study the equations involving boundary values)

Cellwise computations; formulas

- Repeat the previous example, but apply the cellwise algorithm
- Work with one cell at a time
- Transform physical cell to reference cell $X \in [-1,1]$

$$A_{i-1,j-1}^{(e)} = \int_{\Omega^{(e)}} \varphi_i'(x) \varphi_j'(x) \,\mathrm{d}x = \int_{-1}^1 \frac{d}{dx} \tilde{\varphi}_r(X) \frac{d}{dx} \tilde{\varphi}_s(X) \frac{h}{2} \,\mathrm{d}X,$$

$$\tilde{\varphi}_0(X) = \frac{1}{2}(1-X), \quad \tilde{\varphi}_1(X) = \frac{1}{2}(1+X)$$

$$\frac{d\tilde{\varphi}_0}{dX} = -\frac{1}{2}, \quad \frac{d\tilde{\varphi}_1}{dX} = \frac{1}{2}$$

From the chain rule

$$\frac{d\tilde{\varphi}_r}{dx} = \frac{d\tilde{\varphi}_r}{dX}\frac{dX}{dx} = \frac{2}{h}\frac{d\tilde{\varphi}_r}{dX}$$

Cellwise computations; details

$$A_{i-1,j-1}^{(e)} = \int_{\Omega^{(e)}} \varphi_i'(x) \varphi_j'(x) \, \mathrm{d}x = \int_{-1}^1 \frac{2}{h} \frac{d\tilde{\varphi}_r}{dX} \frac{2}{h} \frac{d\tilde{\varphi}_s}{dX} \frac{h}{2} \, \mathrm{d}X = \tilde{A}_{r,s}^{(e)}$$

$$b_{i-1}^{(\mathbf{e})} = \int_{\Omega^{(\mathbf{e})}} 2\varphi_i(x) \, \mathrm{d}x = \int_{-1}^1 2\tilde{\varphi}_r(X) \frac{h}{2} \, \mathrm{d}X = \tilde{b}_r^{(\mathbf{e})}, \quad i = q(\mathbf{e}, r), \ r = 0, 1$$

Must run through all r,s=0,1 and r=0,1 and compute each entry in the element matrix and vector:

$$\tilde{A}^{(e)} = \frac{1}{h} \begin{pmatrix} 1 & -1 \\ -1 & 1 \end{pmatrix}, \quad \tilde{b}^{(e)} = h \begin{pmatrix} 1 \\ 1 \end{pmatrix}$$

Example:

$$\tilde{A}_{0,1}^{(e)} = \int_{-1}^{1} \frac{2}{h} \frac{d\tilde{\varphi}_{0}}{dX} \frac{2}{h} \frac{d\tilde{\varphi}_{1}}{dX} \frac{h}{2} \, \mathrm{d}X = \frac{2}{h} (-\frac{1}{2}) \frac{2}{h} \frac{1}{2} \frac{h}{2} \int_{-1}^{1} \, \mathrm{d}X = -\frac{1}{h}$$

Cellwise computations; details of boundary cells

- The boundary cells involve only one unknown
- ullet $\Omega^{(0)}$: left node value known, only a contribution from right node
- ullet $\Omega^{(N_c)}$: right node value known, only a contribution from left

For e = 0 and $= N_e$

$$\tilde{A}^{(e)} = \frac{1}{h} \begin{pmatrix} 1 \end{pmatrix}, \quad \tilde{b}^{(e)} = h \begin{pmatrix} 1 \end{pmatrix}$$

Only one degree of freedom ("node") in these cells (r=0 counts the only dof)

Cellwise computations; assembly

4 P1 elements:

vertices =
$$[0, 0.5, 1, 1.5, 2]$$

cells = $[[0, 1], [1, 2], [2, 3], [3, 4]]$
dof.map = $[[0], [0, 1], [1, 2], [2]]$ # only 1 dof in elm 0,

Python code for the assembly algorithm:

```
# ke[e][r,s]: element matrix, be[e][r]: element vector
# k[i,j]: coef[scient matrix, b[i]: right-hand side
for e in range(len(Ae)):
    for r in range(ke[e] shape[0]):
        for s in range(ke[e] shape[1]);
        A[dof_map[e,r],dof_map[e,s]] += ke[e][i,j]
        b[dof_map[e,r]] += be[e][i,j]
```

Result: same linear system as arose from computations in the physical domain

General construction of a boundary function

- Now we address nonzero Dirichlet conditions
- B(x) is not always easy to construct (extend to the interior of Ω), especially not in 2D and 3D
- With finite element φ_i , B(x) can be constructed in a completely general way
- Ib: set of indices with nodes where u is known
- U_i : Dirichlet value of u at node $i, i \in I_b$

$$B(x) = \sum_{j \in I_b} U_j \varphi_j(x)$$

Suppose we have a Dirichlet condition $u(x_k) = U_k$, $k \in I_b$:

$$u(x_k) = \sum_{j \in I_k} U_j \underbrace{\varphi_j(x)}_{\neq 0 \text{ only for } j = k} + \sum_{j \in \mathcal{I}_s} c_j \underbrace{\varphi_{\nu(j)}(x_k)}_{=0, \ k \notin \mathcal{I}_s} = U_k$$

Example with two Dirichlet values; variational formulation

$$-u'' = 2$$
, $u(0) = C$, $u(L) = D$

$$\int_0^L u'v' \, \mathrm{d}x = \int_0^L 2v \, \mathrm{d}x \quad \forall v \in V$$

$$(u', v') = (2, v) \quad \forall v \in V$$

Example with two Dirichlet values; boundary function

$$B(x) = \sum_{j \in I_b} U_j \varphi_j(x)$$

Here $I_b = \{0, N_n\}, \ U_0 = C, \ U_{N_n} = D,$

$$\psi_i = \varphi_{\nu(i)}, \quad \nu(i) = i + 1, \quad i \in \mathcal{I}_s = \{0, \dots, N = N_n - 2\}$$

$$u(x) = C\varphi_0(x) + D\varphi_{N_n}(x) + \sum_{j \in \mathcal{I}_n} c_j \varphi_{\nu(j)}$$

Example with two Dirichlet values; details

Insert $u = B + \sum_{i} c_{i} \psi_{i}$ in variational formulation:

$$(u',v')=(2,v) \quad \Rightarrow \quad (\sum_j c_j\psi'_j,\psi'_i)=(2-B',\psi_i) \quad \forall v \in V$$

$$u(x) = \underbrace{C \cdot \varphi_0 + D\varphi_{N_o}}_{B(x)} + \sum_{j \in \mathcal{I}_s} c_j \varphi_{j+1}$$
$$= C \cdot \varphi_0 + D\varphi_{N_o} + c_0 \varphi_1 + c_1 \varphi_2 + \dots + c_N \varphi_{N_o-1}$$

$$A_{i-1,j-1} = \int_0^L \varphi_i'(x) \varphi_j'(x) dx, \quad b_{i-1} = \int_0^L (f(x) - C \varphi_0'(x) - D \varphi_{N_n}'(x)) \varphi_i(x) dx,$$
 for $i, j = 1, \dots, N + 1 = N_n - 1$.

New boundary terms from $-\int B' \varphi_i \, \mathrm{d} x\colon C/2$ for i=1 and -D/2 for $i=N_n-1$

Example with two Dirichlet values; cellwise computations

- Element matrices as in the previous example (with u=0 on the boundary)
- New element vector in the first and last cell

From the last cell:

$$\tilde{b}_{0}^{(N_{e})} = \int_{-1}^{1} \left(f - D \frac{2}{h} \frac{d\tilde{\varphi}_{1}}{dX} \right) \tilde{\varphi}_{0} \frac{h}{2} dX = \left(\frac{h}{2} (2 - D \frac{2}{h} \frac{1}{2}) \int_{-1}^{1} \tilde{\varphi}_{0} dX = h - D \right)$$

From the first cell:

$$\tilde{b}_0^{(0)} = \int_{-1}^1 \left(f - C \frac{2}{h} \frac{d\tilde{\varphi}_0}{dX} \right) \tilde{\varphi}_1 \frac{h}{2} \, \mathrm{d}X = (\frac{h}{2} (2 + C \frac{2}{h} \frac{1}{2}) \int_{-1}^1 \tilde{\varphi}_1 \, \mathrm{d}X = h + C/2$$

Modification of the linear system; ideas

- Method 1: incorporate Dirichlet values through a B(x) function and demand $\psi_i = 0$ where Dirichlet values apply
- Method 2: drop $B(\mathbf{x})$, drop demands to ψ_i , just assemble as if there were no Dirichlet conditions, and modify the linear system instead

Method 2: always $\psi_i = \varphi_i$ and

$$u(x) = \sum_{j \in \mathcal{I}_s} c_j \varphi_j(x), \quad \mathcal{I}_s = \{0, \dots, N = N_n\}$$

Attractive way of incorporating Dirichlet conditions

 \boldsymbol{u} is treated as unknown at all boundaries when computing entries in the linear system

Modification of the linear system; original system

$$-u'' = 2$$
, $u(0) = 0$, $u(L) = D$

Assemble as if there were no Dirichlet conditions:

$$\frac{1}{h} \begin{pmatrix} 1 & -1 & 0 & \cdots & \cdots & \cdots & \cdots & 0 \\ -1 & 2 & -1 & \ddots & & & & & \vdots \\ 0 & -1 & 2 & -1 & \ddots & & & & \vdots \\ \vdots & \ddots & \ddots & \ddots & \ddots & \ddots & & \vdots \\ \vdots & & & 0 & -1 & 2 & -1 & \ddots & \vdots \\ \vdots & & & & \ddots & \ddots & \ddots & \ddots & 0 \\ \vdots & & & & & \ddots & \ddots & \ddots & \ddots & 0 \\ \vdots & & & & & & \ddots & \ddots & \ddots & -1 \\ 0 & \cdots & \cdots & \cdots & \cdots & 0 & -1 & 1 \end{pmatrix} \begin{pmatrix} c_0 \\ \vdots \\ \vdots \\ \vdots \\ \vdots \\ \vdots \\ c_N \end{pmatrix} = \begin{pmatrix} h \\ 2h \\ \vdots \\ \vdots \\ \vdots \\ \vdots \\ c_N \end{pmatrix}$$

Modification of the linear system; row replacement

- ullet Dirichlet condition $u(x_k)=U_k$ means $c_k=U_k$ (since $c_k=u(x_k)$)
- Replace first row by $c_0 = 0$
- Replace last row by $c_N = D$

Modification of the linear system; element matrix/vector

In cell 0 we know u for local node (degree of freedom) r=0. Replace the first cell equation by $\tilde{c}_0=0$:

$$\tilde{A}^{(0)} = A = \frac{1}{h} \begin{pmatrix} h & 0 \\ -1 & 1 \end{pmatrix}, \quad \tilde{b}^{(0)} = \begin{pmatrix} 0 \\ h \end{pmatrix}$$

In cell N_e we know u for local node r=1. Replace the last equation in the cell system by $\tilde{c}_1=D$:

$$\tilde{A}^{(N_e)} = A = \frac{1}{h} \begin{pmatrix} 1 & -1 \\ 0 & h \end{pmatrix}, \quad \tilde{b}^{(N_e)} = \begin{pmatrix} h \\ D \end{pmatrix}$$

Symmetric modification of the linear system; algorithm

- ullet The modification above destroys symmetry of the matrix: e.g., $A_{0,1}
 eq A_{1,0}$
- Symmetry is often important in 2D and 3D (faster computations)
- A more complex modification can preserve symmetry!

Algorithm for incorporating $c_i = U_i$ in a symmetric way:

- \bigcirc Subtract column i times U_i from the right-hand side
- Zero out column and row no i
- Place 1 on the diagonal
- \bigcirc Set $b_i = U_i$

Symmetric modification of the linear system; example

$$\frac{1}{h} \begin{pmatrix} 1 & 0 & 0 & \cdots & \cdots & \cdots & \cdots & 0 \\ 0 & 2 & -1 & \ddots & & & & & \vdots \\ 0 & -1 & 2 & -1 & \ddots & & & & \vdots \\ \vdots & \ddots & \ddots & \ddots & \ddots & \ddots & \vdots \\ \vdots & & & \ddots & \ddots & \ddots & \ddots & \vdots \\ \vdots & & & & 0 & -1 & 2 & -1 & \ddots & \vdots \\ \vdots & & & & & \ddots & \ddots & \ddots & 0 \\ \vdots & & & & & \ddots & \ddots & \ddots & 0 \\ \vdots & & & & & & \ddots & \ddots & 0 \\ \vdots & & & & & & \ddots & \ddots & 0 \\ \vdots & & & & & & \ddots & \ddots & 0 \\ \vdots & & & & & & \ddots & \ddots & 0 \\ \vdots & & & & & & \ddots & \ddots & 0 \\ \vdots & & & & & & & \ddots & 0 \\ \vdots & & & & & & \ddots & 0 \\ \vdots & & & & & & \ddots & 0 \\ \vdots & & & & & & \ddots & \ddots & 0 \\ \vdots & & & & & & \ddots & \ddots & 0 \\ \vdots & & & & & & \ddots & 0 \\ \vdots & & & & & & \ddots & 0 \\ \vdots & & & & & & \ddots & 0 \\ \vdots & &$$

Symmetric modification of the linear system; element level

Symmetric modification applied to $\tilde{A}^{(N_e)}$:

$$\tilde{A}^{(N_e)} = A = \frac{1}{h} \begin{pmatrix} 1 & 0 \\ 0 & 1 \end{pmatrix}, \quad \tilde{b}^{(N-1)} = \begin{pmatrix} h + D/h \\ D \end{pmatrix}$$

Boundary conditions: specified derivative

Neumann conditions

How can we incorporate u'(0) = C with finite elements?

$$-u'' = f$$
, $u'(0) = C$, $u(L) = D$

- $\psi_i(L) = 0$ because of Dirichlet condition u(L) = D
- No demand to $\psi_i(0)$

The variational formulation

Galerkin's method:

$$\int_0^L (u''(x) + f(x))\psi_i(x)dx = 0, \quad i \in \mathcal{I}_s$$

Integration of $u''\psi_i$ by parts:

$$\int_{0}^{L} u'(x)\psi'_{i}(x) dx - (u'(L)\psi_{i}(L) - u'(0)\psi_{i}(0)) - \int_{0}^{L} f(x)\psi_{i}(x) dx = 0, \quad i \in$$

- $u'(L)\psi_i(L) = 0$ since $\psi_i(L) = 0$
- $u'(0)\psi_i(0) = C\psi_i(0)$ since u'(0) = C

Method 1: Boundary function and exclusion of Dirichlet degrees of freedom

- $\psi_i = \varphi_i, i \in \mathcal{I}_s = \{0, ..., N = N_n 1\}$
- $B(x) = D\varphi_{N_n}(x)$, $u = B + \sum_{j=0}^{N} c_j \varphi_j$

$$\int_0^L u'(x)\varphi_i'(x)dx = \int_0^L f(x)\varphi_i(x)dx - C\varphi_i(0), \quad i \in \mathcal{I}_s$$

$$\sum_{i=0}^{N=N_n-1} \left(\int_0^L \varphi_i'(x) \varphi_j'(x) dx \right) c_j = \int_0^L \left(f(x) \varphi_i(x) - D \varphi_N'(x) \varphi_i(x) \right) dx - C$$

for $i = 0, ..., N = N_n - 1$.

Method 2: Use all φ_i and insert the Dirichlet condition in the linear system

- Now $\psi_i = \varphi_i$, $i = 0, ..., N = N_n$
- $\varphi_N(L) \neq 0$, so $u'(L)\varphi_N(L) \neq 0$
- However, the term $u'(L)\varphi_N(L)$ in b_N will be erased when we insert the Dirichlet value in $b_N=D$

We can forget about the term $u'(L)\varphi_i(L)!$

Result

Boundary terms $u'\varphi_i$ at points x_i where Dirichlet values apply can always be forgotten.

$$u(x) = \sum_{j=0}^{N=N_n} c_j \varphi_j(x)$$

$$\sum_{i=N_n}^{N=N_n} \left(\int_{-i}^{L} \varphi_i'(x) \varphi_i'(x) dx \right) c_j = \int_{-i}^{L} f(x) \varphi_i(x) \varphi_i(x) dx - C \varphi_i(0)$$

How the Neumann condition impacts the element matrix and vector

The extra term $C\varphi_0(0)$ affects only the element vector from the first cells since $\varphi_0=0$ on all other cells.

$$\tilde{A}^{(0)} = A = \frac{1}{h} \begin{pmatrix} 1 & 1 \\ -1 & 1 \end{pmatrix}, \quad \tilde{b}^{(0)} = \begin{pmatrix} h - C \\ h \end{pmatrix}$$

The finite element algorithm

The differential equation problem defines the integrals in the variational formulation.

Request these functions from the user:

integrand_lhs(phi, r, s, x)
boundary_lhs(phi, r, s, x)
integrand_rhs(phi, r, x)
boundary_rhs(phi, r, x)

Must also have a mesh with vertices, cells, and dof_map

Python pseudo code; the element matrix and vector

Python pseudo code; boundary conditions and assembly

Variational formulations in 2D and 3D

How to do integration by parts is the major difference when moving to 2D and 3D.

Integration by parts

Rule for multi-dimensional integration by parts

$$-\int_{\Omega} \nabla \cdot (a(x)\nabla u) v \, \mathrm{d}x = \int_{\Omega} a(x)\nabla u \cdot \nabla v \, \mathrm{d}x - \int_{\partial \Omega} a \frac{\partial u}{\partial n} v \, \mathrm{d}s$$

- \int_{Ω} () dx: area (2D) or volume (3D) integral
- $\int_{\partial\Omega}$ () ds: line(2D) or surface (3D) integral
- $\partial \Omega_N$: Neumann conditions $-a \frac{\partial u}{\partial n} = g$
- $\partial\Omega_D$: Dirichlet conditions $u=u_0$
- $v \in V$ must vanish on $\partial \Omega_D$ (in method 1)

Example on integration by parts; problem

$$\begin{aligned} \mathbf{v} \cdot \nabla u + \alpha u &= \nabla \cdot (a \nabla u) + f, & \mathbf{x} \in \Omega \\ u &= u_0, & \mathbf{x} \in \partial \Omega_D \\ -a \frac{\partial u}{\partial x} &= g, & \mathbf{x} \in \partial \Omega_N \end{aligned}$$

- Known: a, α , f, u_0 , and g.
- Second-order PDE: must have exactly one boundary condition at each point of the boundary

Method 1 with boundary function and $\psi_i = 0$ on $\partial \Omega_D$:

$$u(\mathbf{x}) = B(\mathbf{x}) + \sum_{j \in \mathcal{I}_s} c_j \psi_j(\mathbf{x}), \quad B(\mathbf{x}) = u_0(\mathbf{x})$$

Example on integration by parts; details (1)

Galerkin's method: multiply by $v \in V$ and integrate over Ω ,

$$\int_{\Omega} (\boldsymbol{v} \cdot \nabla \boldsymbol{u} + \alpha \boldsymbol{u}) \boldsymbol{v} \, \mathrm{d} \boldsymbol{x} = \int_{\Omega} \nabla \cdot (\boldsymbol{a} \nabla \boldsymbol{u}) \, \, \mathrm{d} \boldsymbol{x} + \int_{\Omega} \boldsymbol{f} \boldsymbol{v} \, \mathrm{d} \boldsymbol{x}$$

Integrate second-order term by parts:

$$\int_{\Omega} \nabla \cdot (a \nabla u) v \, dx = -\int_{\Omega} a \nabla u \cdot \nabla v \, dx + \int_{\partial \Omega} a \frac{\partial u}{\partial n} v \, ds,$$

Resulting variational form:

$$\int_{\Omega} (\mathbf{v} \cdot \nabla \mathbf{u} + \alpha \mathbf{u}) \mathbf{v} \, d\mathbf{x} = -\int_{\Omega} \mathbf{a} \nabla \mathbf{u} \cdot \nabla \mathbf{v} \, d\mathbf{x} + \int_{\partial \Omega} \mathbf{a} \frac{\partial \mathbf{u}}{\partial \mathbf{n}} \mathbf{v} \, d\mathbf{s} + \int_{\Omega} \mathbf{f} \mathbf{v} \, d\mathbf{x}$$

Example on integration by parts; details (2)

Note: $v \neq 0$ only on $\partial \Omega_N$:

$$\int_{\partial\Omega} a \frac{\partial u}{\partial n} v \, \mathrm{d}s = \int_{\partial\Omega_N} \underbrace{a \frac{\partial u}{\partial n}}_{-\sigma} v \, \mathrm{d}s = -\int_{\partial\Omega_N} g v \, \mathrm{d}s$$

The final variational form:

$$\int_{\Omega} (\mathbf{v} \cdot \nabla \mathbf{u} + \alpha \mathbf{u}) \mathbf{v} \, \mathrm{d}\mathbf{x} = -\int_{\Omega} \mathbf{a} \nabla \mathbf{u} \cdot \nabla \mathbf{v} \, \mathrm{d}\mathbf{x} - \int_{\partial \Omega_N} \mathbf{g} \mathbf{v} \, \mathrm{d}\mathbf{s} + \int_{\Omega} \mathbf{f} \mathbf{v} \, \mathrm{d}\mathbf{x}$$

Or with inner product notation:

$$(\mathbf{v} \cdot \nabla \mathbf{u}, \mathbf{v}) + (\alpha \mathbf{u}, \mathbf{v}) = -(\mathbf{a} \nabla \mathbf{u}, \nabla \mathbf{v}) - (\mathbf{g}, \mathbf{v})_N + (\mathbf{f}, \mathbf{v})$$

 $(g,v)_N$: line or surface integral over $\partial\Omega_N$

Example on integration by parts; linear system

$$u = B + \sum_{j \in \mathcal{I}_s} c_j \psi_j, \quad B = u_0$$

$$A_{i,j} = (\mathbf{v} \cdot \nabla \psi_j, \psi_i) + (\alpha \psi_j, \psi_i) + (\mathbf{a} \nabla \psi_j, \nabla \psi_i)$$

$$b_i = (g, \psi_i)_N + (f, \psi_i) - (\mathbf{v} \cdot \nabla u_0, \psi_i) + (\alpha u_0, \psi_i) + (a \nabla u_0, \nabla \psi_i)$$

Transformation to a reference cell in 2D/3D (1)

We want to compute an integral in the physical domain by integrating over the reference cell.

$$\int_{\Omega^{(e)}} a(x) \nabla \varphi_i \cdot \nabla \varphi_j \, dx$$

Mapping from reference to physical coordinates:

with Jacobian J.

$$J_{i,j} = \frac{\partial x_j}{\partial X_i}$$

- $dx \rightarrow \det J dX$.
- Must express $\nabla \varphi_i$ by an expression with $\tilde{\varphi}_r$, i = q(e,r):

Transformation to a reference cell in 2D/3D (2)

Can derive

$$\nabla_{\mathbf{X}}\tilde{\varphi}_{r} = J \cdot \nabla_{\mathbf{X}}\varphi_{i}$$

$$\nabla_{\mathbf{X}}\varphi_{i} = \nabla_{\mathbf{X}}\tilde{\varphi}_{r}(\mathbf{X}) = J^{-1} \cdot \nabla_{\mathbf{X}}\tilde{\varphi}_{r}(\mathbf{X})$$

Integral transformation from physical to reference coordinates:

$$\int_{\Omega^{(s)}} a(\mathbf{x}) \nabla_{\mathbf{x}} \varphi_{i} \cdot \nabla_{\mathbf{x}} \varphi_{j} \, \mathrm{d}\mathbf{x} = \int_{\tilde{\Omega}'} a(\mathbf{x}(\mathbf{X})) (J^{-1} \cdot \nabla_{\mathbf{X}} \tilde{\varphi}_{r}) \cdot (J^{-1} \cdot \nabla \tilde{\varphi}_{s}) \, \mathrm{det} \, J \, \mathrm{d}\mathbf{X}$$

Numerical integration

Numerical integration over reference cell triangles and tetrahedra:

$$\int_{\tilde{\Omega}^r} g \, \mathrm{d}X = \sum_{j=0}^{n-1} w_j g(\bar{\boldsymbol{X}}_j)$$

Module numint.py contains different rules:

- [0.16666666666666666, 0.1666666666666, 0.166666666666666]
- Triangle: rules with n = 1, 3, 4, 7 integrate exactly polynomials of degree 1, 2, 3, 4, resp.
- Tetrahedron: rules with n = 1, 4, 5, 11 integrate exactly polynomials of degree 1, 2, 3, 4, resp.

Time-dependent problems

- So far: used the finite element framework for discretizing in space
- What about $u_t = u_{xx} + f$?
 - Use finite differences in time to obtain a set of recursive spatial problems
 - Solve the spatial problems by the finite element method

Example: diffusion problem

$$\frac{\partial u}{\partial t} = \alpha \nabla^2 u + f(\mathbf{x}, t), \qquad \mathbf{x} \in \Omega, t \in (0, T]$$

$$u(\mathbf{x}, 0) = l(\mathbf{x}), \qquad \mathbf{x} \in \Omega$$

$$\frac{\partial u}{\partial n} = 0, \qquad \mathbf{x} \in \partial \Omega, \ t \in (0, T]$$

A Forward Euler scheme; ideas

$$[D_t^+ u = \alpha \nabla^2 u + f]^n$$
, $n = 1, 2, ..., N_t - 1$

Solving wrt u^{n+1} :

$$u^{n+1} = u^n + \Delta t \left(\alpha \nabla^2 u^n + f(x, t_n) \right)$$

- \bullet $u^n = \sum_j c_j^n \psi_j \in V$, $u^{n+1} = \sum_j c_j^{n+1} \psi_j \in V$
- Compute u^0 from I
- ullet Compute u^{n+1} from u^n by solving the PDE for u^{n+1} at each time level

A Forward Euler scheme; stages in the discretization

- ullet $u_{
 m e}(x,t)$: exact solution of the space-and time-continuous problem
- $u_e^n(x)$: exact solution of time-discrete problem (after applying a finite difference scheme in time)
- $u_n^e(\mathbf{x}) \approx u^n = \sum_{j \in \mathcal{I}_s} c_j^n \psi_j = \text{solution of the time-}$ and space-discrete problem (after applying a Galerkin method in space)

$$\frac{\partial u_{\mathsf{e}}}{\partial t} = \alpha \nabla^2 u_{\mathsf{e}} + f(\mathbf{x}, t)$$

$$u_{\mathsf{e}}^{n+1} = u_{\mathsf{e}}^n + \Delta t \left(\alpha \nabla^2 u_{\mathsf{e}}^n + f(\mathbf{x}, t_n) \right)$$

$$u_{\mathsf{e}}^n \approx u^n = \sum_{j=0}^N c_j^n \psi_j(\mathbf{x}), \quad u_{\mathsf{e}}^{n+1} \approx u^{n+1} = \sum_{j=0}^N c_j^{n+1} \psi_j(\mathbf{x})$$

$$R = u^{n+1} - u^n - \Delta t \left(\alpha \nabla^2 u^n + f(\mathbf{x}, t_n) \right)$$

A Forward Euler scheme; weighted residual (or Galerkin) principle

$$R = u^{n+1} - u^n - \Delta t \left(\alpha \nabla^2 u^n + f(x, t_n) \right)$$

The weighted residual principle:

$$\int_{\Omega} Rw \, \mathrm{d}x = 0, \quad \forall w \in W$$

results in

$$\int_{\Omega} \left[u^{n+1} - u^n - \Delta t \left(\alpha \nabla^2 u^n + f(x, t_n) \right) \right] w \, \mathrm{d}x = 0, \quad \forall w \in W$$

Galerkin: W=V, w=v

A Forward Euler scheme; integration by parts

Isolating the unknown u^{n+1} on the left-hand side:

$$\int_{\Omega} u^{n+1} \psi_i \, \mathrm{d}x = \int_{\Omega} \left[u^n - \Delta t \left(\alpha \nabla^2 u^n + f(x, t_n) \right) \right] v \, \mathrm{d}x$$

Integration by parts of $\int \alpha(\nabla^2 u^n) v \, dx$:

$$\int_{\Omega} \alpha (\nabla^2 u^n) v \, \mathrm{d}x = -\int_{\Omega} \alpha \nabla u^n \cdot \nabla v \, \mathrm{d}x + \underbrace{\int_{\partial \Omega} \alpha \frac{\partial u^n}{\partial n} v \, \mathrm{d}x}_{=0 \quad \leftarrow \quad \partial u^n / \partial n = 0}$$

Variational form:

$$\int_{\Omega} u^{n+1} v \, \mathrm{d}x = \int_{\Omega} u^n v \, \mathrm{d}x - \Delta t \int_{\Omega} \alpha \nabla u^n \cdot \nabla v \, \mathrm{d}x + \Delta t \int_{\Omega} f^n v \, \mathrm{d}x, \quad \forall v \in V$$

New notation for the solution at the most recent time levels

- u and u: the spatial unknown function to be computed
- \bullet $\it u_1$ and u_1: the spatial function at the previous time level $\it t-\Delta\it t$
- u_2 and u_2 : the spatial function at $t-2\Delta t$
- This new notation gives close correspondence between code and math

$$\int_{\Omega} u v \, \mathrm{d}x = \int_{\Omega} u_1 v \, \mathrm{d}x - \Delta t \int_{\Omega} \alpha \nabla u_1 \cdot \nabla v \, \mathrm{d}x + \Delta t \int_{\Omega} f^n v \, \mathrm{d}x$$

or shorter

$$(u, \psi_i) = (u_1, v) - \Delta t(\alpha \nabla u_1, \nabla v) + (f^n, v)$$

Deriving the linear systems

•
$$u = \sum_{j=0}^{N} c_j \psi_j(\mathbf{x})$$

•
$$u_1 = \sum_{j=0}^{N} c_{1,j} \psi_j(x)$$

•
$$\forall v \in V$$
: for $v = \psi_i$, $i = 0, ..., N$

Insert these in

$$(u, \psi_i) = (u_1, \psi_i) - \Delta t(\alpha \nabla u_1, \nabla \psi_i) + (f^n, \psi_i)$$

and order terms as matrix-vector products:

$$\sum_{j=0}^{N} \underbrace{(\psi_i, \psi_j)}_{M_{i,j}} c_j = \sum_{j=0}^{N} \underbrace{(\psi_i, \psi_j)}_{M_{i,j}} c_{1,j} - \Delta t \sum_{j=0}^{N} \underbrace{(\nabla \psi_i, \alpha \nabla \psi_j)}_{K_{i,j}} c_{1,j} + (f^n, \psi_i), \quad i = 0$$

Structure of the linear systems

$$Mc = Mc_1 - \Delta tKc_1 + f$$

$$\begin{aligned} M &= \{M_{i,j}\}, \quad M_{i,j} = (\psi_i, \psi_j), \quad i, j \in \mathcal{I}_s \\ K &= \{K_{i,j}\}, \quad K_{i,j} = (\nabla \psi_i, \alpha \nabla \psi_j), \quad i, j \in \mathcal{I}_s \\ f &= \{(f(\mathbf{x}, t_n), \psi_i)\}_{i \in \mathcal{I}_s} \\ c &= \{c_i\}_{i \in \mathcal{I}_s} \\ c_1 &= \{c_{i,i}\}_{i \in \mathcal{I}_s} \end{aligned}$$

Computational algorithm

- Compute M and K.
- \bigcirc Initialize u^0 by either interpolation or projection
- For $n = 1, 2, ..., N_t$:
 - compute $b = Mc_1 \Delta t Kc_1 + f$
 - solve Mc = b
 - set c₁ = c

Initial condition:

- Either interpolation: $c_{1,j} = I(x_i)$ (finite elements)
- ullet Or projection: solve $\sum_{i} M_{i,j} c_{1,j} = (I,\psi_i), \ i \in \mathcal{I}_{\mathbf{s}}$

Comparing P1 elements with the finite difference method; ideas

- P1 elements in 1D
- Uniform mesh on [0, L] with cell length h
- No Dirichlet conditions: $\psi_i = \varphi_i$, $i = 0, \dots, N = N_n$
- Have found formulas for M and K at the element level
- Have assembled the global matrices
- Have developed corresponding finite difference operator formulas
- $M: h[D_t^+(u+\frac{1}{6}h^2D_xD_xu)]_i^n$
- $K: h[\alpha D_x D_x u]_i^n$

Comparing P1 elements with the finite difference method; results

Diffusion equation with finite elements is equivalent to

$$[D_t^+(u+\frac{1}{6}h^2D_xD_xu)=\alpha D_xD_xu+f]_i^n$$

Can lump the mass matrix by Trapezoidal integration and get the standard finite difference scheme

$$[D_t^+ u = \alpha D_x D_x u + f]_i^n$$

Discretization in time by a Backward Euler scheme

Backward Euler scheme in time:

$$[D_t^- u = \alpha \nabla^2 u + f(x, t)]^n$$

$$u_{\mathsf{e}}^{n} - \Delta t \left(\alpha \nabla^{2} u_{\mathsf{e}}^{n} + f(\mathbf{x}, t_{n}) \right) = u_{\mathsf{e}}^{n-1}$$

$$u_{\mathsf{e}}^{n} \approx u^{n} = \sum_{j=0}^{N} c_{j}^{n} \psi_{j}(\mathbf{x}), \quad u_{\mathsf{e}}^{n+1} \approx u^{n+1} = \sum_{j=0}^{N} c_{j}^{n+1} \psi_{j}(\mathbf{x})$$

The variational form of the time-discrete problem

$$\int_{\Omega} \left(u^n v + \Delta t \alpha \nabla u^n \cdot \nabla v \right) \, \mathrm{d}x = \int_{\Omega} u^{n-1} v \, \mathrm{d}x - \Delta t \int_{\Omega} f^n v \, \mathrm{d}x, \quad \forall v \in V$$
 ...

$$(u, v) + \Delta t(\alpha \nabla u, \nabla v) = (u_1, v) + \Delta t(f^n, \psi_i)$$

The linear system: insert $u=\sum_i c_j \psi_i$ and $u_1=\sum_i c_{1,j} \psi_i$,

$$(M + \Delta t \alpha K)c = Mc_1 + f$$

Calculations with P1 elements in 1D

Can interpret the resulting equation system as

$$[D_t^-(u+\frac{1}{6}h^2D_xD_xu)=\alpha D_xD_xu+f]_i^n$$

Lumped mass matrix (by Trapezoidal integration) gives a standard finite difference method:

$$[D_t^- u = \alpha D_x D_x u + f]_i^n$$

Dirichlet boundary conditions

Dirichlet condition at x = 0 and Neumann condition at x = L:

$$u(x,t) = u_0(x,t),$$
 $x \in \partial \Omega_D$ $-\alpha \frac{\partial}{\partial x} u(x,t) = g(x,t),$ $x \in \partial \Omega_N$

Forward Euler in time, Galerkin's method, and integration by parts:

$$\int_{\Omega} u^{n+1} v \, \mathrm{d}x = \int_{\Omega} (u^n - \Delta t \alpha \nabla u^n \cdot \nabla v) \, \mathrm{d}x - \Delta t \int_{\partial \Omega_N} g v \, \mathrm{d}s, \quad \forall v \in V$$

Requirement: v=0 on $\partial\Omega_D$

Boundary function

$$u^n(\mathbf{x}) = u_0(\mathbf{x}, t_n) + \sum_{j \in \mathcal{I}_s} c_j^n \psi_j(\mathbf{x})$$

$$\begin{split} \sum_{j \in \mathcal{I}_s} \left(\int_{\Omega} \psi_i \psi_j \, \mathrm{d}x \right) c_j^{n+1} &= \sum_{j \in \mathcal{I}_s} \left(\int_{\Omega} \left(\psi_i \psi_j - \Delta t \alpha \nabla \psi_i \cdot \nabla \psi_j \right) \, \mathrm{d}x \right) c_j^{n} - \\ & \int_{\Omega} \left(u_0(\mathbf{x}, t_{n+1}) - u_0(\mathbf{x}, t_n) + \Delta t \alpha \nabla u_0(\mathbf{x}, t_n) + \Delta t \alpha \nabla u_0(\mathbf{x}, t_n) \right) \\ & + \Delta t \int_{\Omega} f \psi_i \, \mathrm{d}x - \Delta t \int_{\partial \Omega_N} g \psi_i \, \mathrm{d}s, \quad i \in \mathcal{I}_s \end{split}$$

Finite element basis functions

- $\begin{array}{l} \bullet \; B(\mathbf{x},t_n) = \sum_{j \in I_b} U_j^n \varphi_j \\ \bullet \; \; \psi_i = \varphi_{\nu(j)}, \; j \in \mathcal{I}_s \\ \bullet \; \; \nu(j), \; j \in \mathcal{I}_s, \; \text{are the node numbers corresponding to all nodes} \end{array}$ without a Dirichlet condition

$$\begin{split} u^n &= \sum_{j \in I_b} U_j^n \varphi_j + \sum_{j \in \mathcal{I}_s} c_{1,j} \varphi_{\nu(j)}, \\ u^{n+1} &= \sum_{j \in I_b} U_j^{n+1} \varphi_j + \sum_{j \in \mathcal{I}_s} c_{j} \varphi_{\nu(j)} \end{split}$$

$$\begin{split} \sum_{j \in \mathcal{I}_s} \left(\int_{\Omega} \varphi_i \varphi_j \, \mathrm{d} \mathbf{x} \right) c_j &= \sum_{j \in \mathcal{I}_s} \left(\int_{\Omega} \left(\varphi_i \varphi_j - \Delta t \alpha \nabla \varphi_i \cdot \nabla \varphi_j \right) \, \mathrm{d} \mathbf{x} \right) c_{1,j} - \\ & \sum_{j \in I_b} \int_{\Omega} \left(\varphi_i \varphi_j (U_j^{n+1} - U_j^n) + \Delta t \alpha \nabla \varphi_i \cdot \nabla \varphi_j U_j^n \right) \, \mathrm{d} \mathbf{x} \\ &+ \Delta t \int_{\Omega} f \varphi_i \, \mathrm{d} \mathbf{x} - \Delta t \int_{\partial \Omega_N} g \varphi_i \, \mathrm{d} \mathbf{s}, \quad i \in \mathcal{I}_s \end{split}$$

Modification of the linear system; the raw system

- Drop boundary function
- Compute as if there are not Dirichlet conditions
- Modify the linear system to incorporate Dirichlet conditions
- \mathcal{I}_s holds the indices of all nodes $\{0, 1, \dots, N = N_n\}$

$$\sum_{j \in \mathcal{I}_{s}} \left(\underbrace{\int_{\Omega} \varphi_{i} \varphi_{j} \, \mathrm{d}x}_{M_{i,j}} \right) c_{j} = \sum_{j \in \mathcal{I}_{s}} \left(\underbrace{\int_{\Omega} \varphi_{i} \varphi_{j} \, \mathrm{d}x}_{M_{i,j}} - \Delta t \underbrace{\int_{\Omega} \alpha \nabla \varphi_{i} \cdot \nabla \varphi_{j} \, \mathrm{d}x}_{K_{i,j}} \right) c_{1,j}$$

$$\underbrace{-\Delta t \int_{\Omega} f \varphi_{i} \, \mathrm{d}x - \Delta t \int_{\partial \Omega_{N}} g \varphi_{i} \, \mathrm{d}s}_{f_{i}}, \quad i \in \mathcal{I}_{s}$$

Modification of the linear system; setting Dirichlet conditions

$$Mc = b$$
, $b = Mc_1 - \Delta t K c_1 + \Delta t f$

For each k where a Dirichlet condition applies, $u(x_k, t_{n+1}) = U_k^{n+1}$,

- \bullet set row k in M to zero and 1 on the diagonal: $M_{k,j}=0,$ $j\in\mathcal{I}_s,\ M_{k,k}=1$
- $b_k = U_k^{n+1}$

Or apply the slightly more complicated modification which preserves symmetry of ${\it M}$

Modification of the linear system; Backward Euler example

Backward Euler discretization in time gives a more complicated coefficient matrix:

$$Ac = b$$
, $A = M + \Delta tK$, $b = Mc_1 + \Delta tf$

- ullet Set row k to zero and 1 on the diagonal: $M_{k,j}=0,\,j\in\mathcal{I}_{\mathbf{s}},$ $M_{k,k}=1$
- Set row k to zero: $K_{k,j} = 0$, $j \in \mathcal{I}_s$
- $b_k = U_k^{n+1}$

Observe: $A_{k,k} = M_{k,k} + \Delta t K_{k,k} = 1 + 0$, so $c_k = U_k^{n+1}$

Analysis of the discrete equations

The diffusion equation $u_t = \alpha u_{\mathrm{xx}}$ allows a (Fourier) wave component

$$u = A_e^n e^{ikx}, A_e = e^{-\alpha k^2 \Delta t}$$

Numerical schemes often allow the similar solution

$$u_n^n = A^n e^{ikx}$$

- A: amplification factor to be computed
- How good is this A compared to the exact one?

Handy formulas

$$\begin{split} & \left[D_t^+ A^n e^{ikq\Delta x}\right]^n = A^n e^{ikq\Delta x} \frac{A-1}{\Delta t}, \\ & \left[D_t^- A^n e^{ikq\Delta x}\right]^n = A^n e^{ikq\Delta x} \frac{1-A^{-1}}{\Delta t}, \\ & \left[D_t A^n e^{ikq\Delta x}\right]^{n+\frac{1}{2}} = A^{n+\frac{1}{2}} e^{ikq\Delta x} \frac{A^{\frac{1}{2}}-A^{-\frac{1}{2}}}{\Delta t} = A^n e^{ikq\Delta x} \frac{A-1}{\Delta t}, \end{split}$$

$$[D_x D_x A^n e^{ikq\Delta x}]_q = -A^n \frac{4}{\Delta x^2} \sin^2 \left(\frac{k\Delta x}{2}\right)$$

Amplification factor for the Forward Euler method; results

Introduce $p=k\Delta x/2$ and $C=\alpha\Delta t/\Delta x^2$:

$$A = 1 - 4C \frac{\sin^2 p}{1 - \frac{2}{3}\sin^2 p}$$
from M

(See notes for details)

Stability: $|A| \leq 1$:

$$C \leq \frac{1}{6} \quad \Rightarrow \quad \Delta t \leq \frac{\Delta x^2}{6\alpha}$$

Finite differences: $C \leq \frac{1}{2}$, so finite elements give a *stricter* stability criterion for this PDE!

Amplification factor for the Backward Euler method; results

Coarse meshes:

$$A = \left(1 + 4C \frac{\sin^2 p}{1 + \frac{2}{3}\sin^2 p}\right)^{-1} \text{ (unconditionally stable)}$$

